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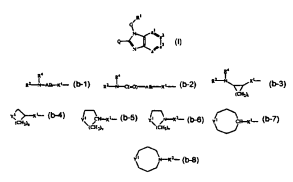
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[Continued on next page]

(54) Title: RESPIRATORY SYNCYTIAL VIRUS REPLICATION INHIBITORS



(57) Abstract: The present invention concerns compounds of formula (I), prodrugs, N-oxides, addition salts, quaternary amines, metal complexes and stereochemically isomeric forms thereof wherein -a1=a2-a3=a4- represents a radical of formula -CH=CH-CH=CH-; -N=CH-CH=CH-; -CH=N-CH=CH-; -CH=CH-N=CH-; -CH=CH-CH=N-; wherein each hydrogen atom may optionally be substituted; Q is a radical of formulae (b-1), (b-2), (b-3), (b-4), (b-5), (b-6), (b-7) and (b-8), wherein Alk is C₁₋₆alkanediyl; Y¹ is a bivalent radical of formula -NR²- or -CH(NR²R⁴)-; X¹ is NR⁴, S, S(=O), S(=O)₂, O, CH₂, C(=O), CH(=CH₂), CH(OH), CH(CH₃), CH(OCH₃), CH(SCH₃), CH(NR^{5a}R^{5b}), CH₂-NR⁴ or NR⁴-CH₂; X² is a direct bond, CH₂, C(=O), NR⁴, C₁₋₄alkyl-NR⁴, NR⁴-C₁₋₄alkyl, t is 2 to 5; u is 1 to 5; v is 2 or 3; and whereby each hydrogen in Alk and in (b-3), (b-4), (b-5), (b-6), (b-7) and (b-8), may optionally be replaced by R3; provided that when R3 is hydroxy or C1-alkyloxy, then R3 cannot replace a hydrogen atom in the α position relative to a nitrogen atom; G is a direct bond or optionally substituted C₁₋₁₀alkanediyl; R¹ is an optionally substituted bicyclic heterocycle; R2 is hydrogen, formyl, C1-salkylcarbonyl, Hetcarbonyl, pyrrolidinyl, piperidinyl, homopiperidinyl, C3.7cycloalkyl or C1.10alkyl substituted with N(R6)2 and optionally with another substituent; R3 is hydrogen, hydroxy, C1-6alkyl, C1-6alkyloxy, arylC1-6alkyl or arylC1-6alkyloxy, R4 is hydrogen, C1-6alkyl or arylC1-6alkyl; R5a, R5b, R5c and R5d are hydrogen or C1-salkyl; or R5a and R5b, or R5c and R5d taken together from a bivalent radical of formula -(CH2)s- wherein S is 4 or 5; R6 is hydrogen, C1.4alkyl, formyl, hydroxyC1.6alkyl, C1.6alkylcarbonyl or C1.6alkyloxycarbonyl; aryl is optionally substituted phenyl; Het is pyridyl, pyrimidinyl, pyryzinyl, pyridazinyl; as respiratory syncytial virus replication inhibitors; their preparation, compositions containing them and their use as a medicine.



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RESPIRATORY SYNCYTIAL VIRUS REPLICATION INHIBITORS

- The present invention is concerned with benzimidazoles and imidazopyridines having antiviral activity, in particular, they have an inhibitory activity on the replication of the respiratory syncytial virus. It further concerns their preparation and compositions comprising them, as well as their use as a medicine.
- Human RSV or Respiratory Syncytial Virus is a large RNA virus, member of the family of Paramyxoviridae, subfamily pneumovirinae together with bovine RSV virus. Human RSV is responsible for a spectrum of respiratory tract diseases in people of all ages throughout the world. It is the major cause of lower respiratory tract illness during infancy and childhood. Over half of all infants encounter RSV in their first year of life, and almost all within their first two years. The infection in young children can cause lung damage that persists for years and may contribute to chronic lung disease in later life (chronic wheezing, asthma). Older children and adults often suffer from a (bad) common cold upon RSV infection. In old age, susceptibility again increases, and RSV has been implicated in a number of outbreaks of pneumonia in the aged resulting in significant mortality.

Infection with a virus from a given subgroup does not protect against a subsequent infection with an RSV isolate from the same subgroup in the following winter season. Re-infection with RSV is thus common, despite the existence of only two subtypes, A and B.

Today only three drugs have been approved for use against RSV infection. Ribavirin, a nucleoside analogue, provides an aerosol treatment for serious RSV infection in hospitalized children. The aerosol route of administration, the toxicity (risk of teratogenicity), the cost and the highly variable efficacy limit its use. The other two drugs, RespiGam® and palivizumab, polyclonal and monoclonal antibody immunostimulants, are intended to be used in a preventive way.

Other attempts to develop a safe and effective RSV vaccine have all met with failure thus far. Inactivated vaccines failed to protect against disease, and in fact in some cases enhanced disease during subsequent infection. Life attenuated vaccines have been tried with limited success. Clearly there is a need for an efficacious non-toxic and easy to administer drug against RSV replication.

EP-A-0,005,318, EP-A-0,099,139, EP-A-0,145,037, EP-A-0,144,101, EP-A-0,151,826, EP-A-0,151,824, EP-A-0,232,937, EP-A-0,295,742, EP 0,297,661, EP-A-0,307,014, WO 92 01697 describe benzimidazole and imidazopyridine substituted piperidine and piperazine derivatives as antihistaminics, antiallergics or serotonine antagonists.

Thus, the present invention concerns the compounds of formula (I)

$$Q = \begin{bmatrix} R^1 \\ N \end{bmatrix} \begin{bmatrix} a^1 \\ a^2 \end{bmatrix}$$
 (I)

their prodrugs, N-oxides, addition salts, quaternary amines, metal complexes and stereochemically isomeric forms wherein

-a¹=a²-a³=a⁴- represents a bivalent radical of formula

-CH=CH-CH=CH-

(a-1);

-N=CH-CH=CH-

(a-2);

-CH=N-CH=CH-

(a-3);

-CH=CH-N=CH-

(a-4); or

-CH=CH-CH=N-

(a-5);

wherein each hydrogen atom in the radicals (a-1), (a-2), (a-3), (a-4) and (a-5) may optionally be replaced by halo, C_{1-6} alkyl, nitro, amino, hydroxy,

 $C_{1\text{-}6}$ alkyloxy, polyhalo $C_{1\text{-}6}$ alkyl, carboxyl, amino $C_{1\text{-}6}$ alkyl, mono- or di $(C_{1\text{-}4}$ alkyl)amino $C_{1\text{-}6}$ alkyl, $C_{1\text{-}6}$ alkyloxycarbonyl, hydroxy $C_{1\text{-}6}$ alkyl, or a radical of formula

wherein =Z is =O, =CH-C(=O)-NR 5a R 5b , =CH $_2$, =CH-C $_{1-6}$ alkyl, =N-OH or =N-O-C $_{1-6}$ alkyl;

Q is a radical of formula

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$$R^{2}$$
 R^{2}
 R^{2

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$$Y_1$$
 CH— X^1 — Y_1 N— X^2 — Y_1 CH— X^1 — Y_1 N— X^2 — (b-5) (b-6) (b-7) (b-8)

wherein Alk is C₁₋₆alkanediyl;

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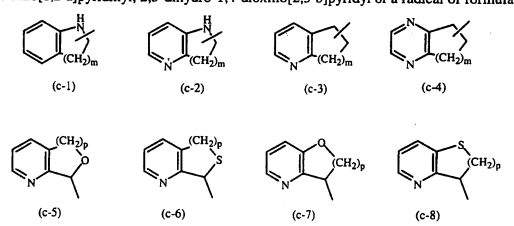
 Y^1 is a bivalent radical of formula $-NR^2$ - or $-CH(NR^2R^4)$ -; X^1 is NR^4 , S, S(=O), S(=O)₂, O, CH₂, C(=O), C(=CH₂), CH(OH), CH(CH₃), CH(OCH₃), CH(SCH₃), CH(NR^{5a}R^{5b}), CH₂-NR⁴ or NR⁴-CH₂; X^2 is a direct bond, CH₂, C(=O), NR⁴, C₁₋₄alkyl-NR⁴, NR⁴-C₁₋₄alkyl; t is 2, 3, 4 or 5; u is 1, 2, 3, 4 or 5; v is 2 or 3; and

whereby each hydrogen atom in Alk and the carbocycles and the heterocycles defined in radicals (b-3), (b-4), (b-5), (b-6), (b-7) and (b-8) may optionally be replaced by R³; with the proviso that when R³ is hydroxy or C₁₋₆alkyloxy, then R³ can not replace a hydrogen atom in the α position relative to a nitrogen atom;

G is a direct bond or C₁₋₁₀alkanediyl optionally substituted with one, two or three substituents selected from hydroxy, C₁₋₆alkyloxy, arylC₁₋₆alkyloxy, C₁₋₆alkylthio, arylC₁₋₆alkylthio, arylcarbonyl, HO(-CH₂-CH₂-O)_n-, C₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, arylC₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, amino, mono-or di(C₁₋₆alkyl)amino, C₁₋₆alkyloxycarbonylamino and aryl;

R¹ is a bicyclic heterocycle selected from quinolinyl, quinoxalinyl, benzofuranyl,

benzothienyl, benzimidazolyl, benzoxazolyl, benzthiazolyl, pyridopyridyl,
naphthiridinyl, 1*H*-imidazo[4,5-b]pyridinyl, 3*H*-imidazo[4,5-b]pyridinyl,
imidazo[1,2-a]pyridinyl, 2,3-dihydro-1,4-dioxino[2,3-b]pyridyl or a radical of formula



and said bicyclic heterocycles may optionally be substituted in either of the two cycles with 1 or where possible more, such as 2, 3 or 4, substituents selected from halo, hydroxy, amino, cyano, carboxy, C₁₋₆alkyl, C₁₋₆alkyloxy, C₁₋₆alkylthio, C₁₋₆alkyloxy-C₁₋₆alkyl, arylC₁₋₆alkyl, arylC₁₋₆alkyloxy, hydroxyC₁₋₆alkyl, mono-or di(C₁₋₆alkyl)-amino, mono-or di(C₁₋₆alkyl)aminoC₁₋₆alkyl, polyhaloC₁₋₆alkyl, C₁₋₆alkylcarbonylamino, C₁₋₆alkyl-SO₂-NR^{5c}-, aryl-SO₂-NR^{5c}-, C₁₋₆alkyloxycarbonyl, -C(=O)-NR^{5c}R^{5d}, HO(-CH₂-CH₂-O)_n-, halo(-CH₂-CH₂-O)_n-, C₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, arylC₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, arylC₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, and mono-or di(C₁₋₆alkyl)amino(-CH₂-CH₂-O)_n-; each n independently is 1, 2, 3 or 4;

each m independently is 1, 2, 3 of 4 each m independently is 1 or 2; each p independently is 1 or 2; each R² independently is hydrogen

each R^2 independently is hydrogen, formyl, C_{1-6} alkylcarbonyl, Hetcarbonyl, pyrrolidinyl, piperidinyl, homopiperidinyl, C_{3-7} cycloalkyl substituted with $N(R^6)_2$, or C_{1-10} alkyl substituted with $N(R^6)_2$ and optionally with a second, third or fourth

substituent selected from amino, hydroxy, C₃₋₇cycloalkyl, C₂₋₅alkanediyl, piperidinyl, mono-or di(C₁₋₆alkyl)amino, C₁₋₆alkyloxycarbonylamino, aryl and aryloxy; R³ is hydrogen, hydroxy, C₁₋₆alkyl, C₁₋₆alkyloxy, arylC₁₋₆alkyl or arylC₁₋₆alkyloxy; R⁴ is hydrogen, C₁₋₆alkyl or arylC₁₋₆alkyl;

 R^{5a} , R^{5b} , R^{5c} and R^{5d} each independently are hydrogen or C_{1-6} alkyl; or

20 R^{5a} and R^{5b}, or R^{5c} and R^{5d} taken together form a bivalent radical of formula -(CH₂)_s-wherein s is 4 or 5;

 R^6 is hydrogen, C_{1-6} alkyl, formyl, hydroxy C_{1-6} alkyl, C_{1-6} alkyloxycarbonyl;

aryl is phenyl or phenyl substituted with 1 or more, such as 2, 3 or 4, substituents selected from halo, hydroxy, C₁₋₆alkyl, hydroxyC₁₋₆alkyl, polyhaloC₁₋₆alkyl, and C₁₋₆alkyloxy;

Het is pyridyl, pyrimidinyl, pyrazinyl, pyridazinyl.

The term prodrug as used throughout this text means the pharmacologically acceptable derivatives, e.g. esters and amides, such that the resulting biotransformation product of the derivative is the active drug as defined in the compounds of formula (I). The reference by Goodman and Gilman (The Pharmacological Basis of Therapeutics, 8th ed., McGraw-Hill, Int. Ed. 1992, "Biotransformation of Drugs", p. 13-15) describing prodrugs generally, is hereby incorporated.

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As used herein C₁₋₃alkyl as a group or part of a group defines straight or branched chain saturated hydrocarbon radicals having from 1 to 3 carbon atoms such as methyl, ethyl,

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propyl, 1-methylethyl and the like; C1-alkyl as a group or part of a group defines straight or branched chain saturated hydrocarbon radicals having from 1 to 4 carbon atoms such as the group defined for C₁₋₃alkyl and butyl and the like; C₂₋₄alkyl as a group or part of a group defines straight or branched chain saturated hydrocarbon radicals having from 2 to 4 carbon atoms such as ethyl, propyl, 1-methylethyl, butyl and 5 the like; C₁₋₆alkyl as a group or part of a group defines straight or branched chain saturated hydrocarbon radicals having from 1 to 6 carbon atoms such as the groups defined for C₁₋₄alkyl and pentyl, hexyl, 2-methylbutyl and the like; C₁₋₉alkyl as a group or part of a group defines straight or branched chain saturated hydrocarbon radicals 10 having from 1 to 9 carbon atoms such as the groups defined for C₁₋₆alkyl and heptyl, octyl, nonyl, 2-methylhexyl, 2-methylheptyl and the like; C₁₋₁₀alkyl as a group or part of a group defines straight or branched chain saturated hydrocarbon radicals having from 1 to 10 carbon atoms such as the groups defined for C_{1.9}alkyl and decyl, 2-methylnonyl and the like. C3-7cycloalkyl is generic to cyclopropyl, cyclobutyl, cyclopentyl, 15 cyclohexyl and cycloheptyl; C_{2.5}alkanediyl defines bivalent straight and branched chain saturated hydrocarbon radicals having from 2 to 5 carbon atoms such as, for example, 1,2-ethanediyl, 1,3-propanediyl, 1,4-butanediyl, 1,2-propanediyl, 2,3-butanediyl, 1,5pentanediyl and the like, C_{2-5} alkanediyl is substituted on C_{1-10} alkyl as provided for in the definition of R², it is meant to be substituted on one carbon atom thus forming a 20 spiro moiety; C₁₋₄alkanediyl defines bivalent straight and branched chain saturated hydrocarbon radicals having from 1 to 4 carbon atoms such as, for example, methylene, 1,2-ethanediyl, 1,3-propanediyl, 1,4-butanediyl and the like; C_{1-6} alkanediyl is meant to include C₁₋₄alkanediyl and the higher homologues thereof having from 5 to 6 carbon atoms such as, for example, 1,5-pentanediyl, 1,6-hexanediyl and the like; 25 C₁₋₁₀alkanediyl is meant to include C₁₋₆alkanediyl and the higher homologues thereof having from 7 to 10 carbon atoms such as, for example, 1,7-heptanediyl, 1,8-octanediyl, 1,9-nonanediyl, 1,10-decanediyl and the like.

As used herein before, the term (=O) forms a carbonyl moiety when attached to a carbon atom, a sulfoxide moiety when attached to a sulfur atom and a sulfonyl moiety when two of said terms are attached to a sulfur atom. The term (=N-OH) forms a hydroxylimine moiety when attached to a carbon atom.

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The term halo is generic to fluoro, chloro, bromo and iodo. As used in the foregoing and hereinafter, polyhaloC₁₋₆alkyl as a group or part of a group is defined as mono- or polyhalosubstituted C₁₋₆alkyl, in particular methyl with one or more fluoro atoms, for example, difluoromethyl or trifluoromethyl. In case more than one halogen atoms are

attached to an alkyl group within the definition of polyhaloC₁₋₄alkyl, they may be the same or different.

As described hereinabove, R¹ defines a bicyclic heterocycle which may optionally be substituted. The substituents may be divided over both rings or they may be attached to one and the same ring.

When any variable (e.g. aryl, R², R³, R⁴, R^{5a}, R^{5b} etc.) occurs more than one time in any constituent, each definition is independent.

It will be appreciated that some of the compounds of formula (I) and their prodrugs, N-oxides, addition salts, quaternary amines, metal complexes and stereochemically isomeric forms may contain one or more centers of chirality and exist as stereochemically isomeric forms.

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The term "stereochemically isomeric forms" as used hereinbefore defines all the possible stereoisomeric forms which the compounds of formula (I), and their prodrugs, N-oxides, addition salts, quaternary amines or physiologically functional derivatives may possess. Unless otherwise mentioned or indicated, the chemical designation of compounds denotes the mixture of all possible stereochemically isomeric forms, said mixtures containing all diastereomers and enantiomers of the basic molecular structure as well as each of the individual isomeric forms of formula (I) and their prodrugs, N-oxides, salts, solvates or quaternary amines substantially free, i.e. associated with less than 10%, preferably less than 5%, in particular less than 2% and most preferably less than 1% of the other isomers. Stereochemically isomeric forms of the compounds of formula (I) are obviously intended to be embraced within the scope of this invention. As used hereinafter the terms trans or cis are well-known by the person skilled in the art.

For therapeutic use, salts of the compounds of formula (I) are those wherein the counterion is pharmaceutically acceptable. However, salts of acids and bases which are non-pharmaceutically acceptable may also find use, for example, in the preparation or purification of a pharmaceutically acceptable compound. All salts, whether pharma-

ceutically acceptable or not are included within the ambit of the present invention.

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The pharmaceutically acceptable acid and base addition salts as mentioned hereinabove are meant to comprise the therapeutically active non-toxic acid and base addition salt forms which the compounds of formula (I) are able to form. The pharmaceutically

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acceptable acid addition salts can conveniently be obtained by treating the base form with such appropriate acid. Appropriate acids comprise, for example, inorganic acids such as hydrohalic acids, e.g. hydrochloric or hydrobromic acid, sulfuric, nitric, phosphoric and the like acids; or organic acids such as, for example, acetic, propanoic, hydroxyacetic, lactic, pyruvic, oxalic (i.e. ethanedioic), malonic, succinic (i.e. butanedioic acid), maleic, fumaric, malic (i.e. hydroxybutanedioic acid), tartaric, citric, methanesulfonic, ethanesulfonic, benzenesulfonic, p-toluenesulfonic, cyclamic, salicylic, p-aminosalicylic, pamoic and the like acids.

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10 Conversely said salt forms can be converted by treatment with an appropriate base into the free base form.

The compounds of formula (I) containing an acidic proton may also be converted into their non-toxic metal or amine addition salt forms by treatment with appropriate organic and inorganic bases. Appropriate base salt forms comprise, for example, the ammonium salts, the alkali and earth alkaline metal salts, e.g. the lithium, sodium, potassium, magnesium, calcium salts and the like, salts with organic bases, e.g. the benzathine, N-methyl-D-glucamine, hydrabamine salts, and salts with amino acids such as, for example, arginine, lysine and the like.

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The term addition salt as used hereinabove also comprises the solvates which the compounds of formula (I) as well as the salts thereof, are able to form. Such solvates are for example hydrates, alcoholates and the like.

The term "quaternary amine" as used hereinbefore defines the quaternary ammonium salts which the compounds of formula (I) are able to form by reaction between a basic nitrogen of a compound of formula (I) and an appropriate quaternizing agent, such as, for example, an optionally substituted alkylhalide, arylhalide or arylalkylhalide, e.g. methyliodide or benzyliodide. Other reactants with good leaving groups may also be used, such as alkyl trifluoromethanesulfonates, alkyl methanesulfonates, and alkyl ptoluenesulfonates. A quaternary amine has a positively charged nitrogen. Pharmaceutically acceptable counterions include chloro, bromo, iodo, trifluoroacetate and acetate. The counterion of choice can be introduced using ion exchange resins.

It will be appreciated that the compounds of formula (I) may have metal binding, chelating, complexating properties and therefore may exist as metal complexes or metal chelates. Such metalated derivatives of the compounds of formula (I) are intended to be included within the scope of the present invention.

Some of the compounds of formula (I) may also exist in their tautomeric form. Such forms although not explicitly indicated in the above formula are intended to be included within the scope of the present invention.

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A special group of com pounds are those compounds of formula (I) wherein one or more of the following restrictions apply:

- Q is a radical of formula (b-1), (b-3), (b-4), (b-5), (b-6), (b-7) or (b-8);
- X² is a direct bond, CH₂ or C(=O);
- R¹ is a bicyclic heterocycle selected from quinolinyl, quinoxalinyl, benzofuranyl, benzothienyl, benzimidazolyl, benzoxazolyl, benzthiazolyl, pyridopyridyl, naphthiridinyl, 1*H*-imidazo[4,5-b]pyridinyl, 3*H*-imidazo[4,5-b]pyridinyl, imidazo[1,2-a]pyridinyl, or a radical of formula

$$(CH_{2})_{m} \qquad (CH_{2})_{m} \qquad (CH_{2})_{p} \qquad (CH_$$

- and said bicyclic heterocycles may optionally be substituted in either of the two cycles with 1 or where possible more, such as 2, 3 or 4, substituents selected from halo, hydroxy, amino, cyano, carboxy, C₁₋₆alkyl, C₁₋₆alkyloxy, C₁₋₆alkylthio, C₁₋₆alkyloxy-C₁₋₆alkyl, arylC₁₋₆alkyl, arylC₁₋₆alkyloxy, hydroxyC₁₋₆alkyl, mono-or di(C₁₋₆alkyl)-amino, mono-or di(C₁₋₆alkyl)aminoC₁₋₆alkyl, polyhaloC₁₋₆alkyl, C₁₋₆alkylcarbonylamino,
- C₁₋₆alkyl-SO₂-NR^{5c}-, aryl-SO₂-NR^{5c}-, C₁₋₆alkyloxycarbonyl, -C(=O)-NR^{5c}R^{5d}, HO(-CH₂-CH₂-O)_n-, halo(-CH₂-CH₂-O)_n-, C₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, arylC₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, arylC₁₋₆alkyloxy(-CH₂-CH₂-O)_n-; each n independently is 1, 2, 3 or 4; each m independently is 1 or 2;
- each p independently is 1 or 2;
 - each R^2 independently is hydrogen, pyrrolidinyl, piperidinyl, homopiperidinyl, C_{3-7} cycloalkyl substituted with NHR⁶, or C_{1-10} alkyl substituted with NHR⁶ and optionally with a second, third or fourth substituent selected from amino, hydroxy,

 C_{3-7} cycloalkyl, C_{2-5} alkanediyl, piperidinyl, mono-or di(C_{1-6} alkyl)amino, C_{1-6} alkyloxycarbonylamino, aryl and aryloxy;

- R³ is hydrogen, hydroxy, C₁₋₆alkyl, C₁₋₆alkyloxy or arylC₁₋₆alkyl;
- R⁶ is hydrogen, C₁₋₆alkyl, formyl, C₁₋₆alkylcarbonyl or C₁₋₆alkyloxycarbonyl.

Another special group of compounds are those compounds wherein $-a^1=a^2-a^3=a^4$ is a radical of formula (a-1), (a-2) or (a-3).

Yet another special group of compounds are those compounds wherein Q is a radical of formula (b-5) wherein v is 2, and Y¹ is -NR²-.

Also interesting compounds are those compounds wherein R^2 is C_{1-10} alkyl substituted with NHR⁶.

- Other interesting compounds are those compounds wherein G is a direct bond or C₁₋₁₀alkanediyl optionally substituted with one two or three substituents selected from hydroxy, C₁₋₆alkyloxy, arylC₁₋₆alkyloxy, HO(-CH₂-CH₂-O)_n-, C₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, arylC₁₋₆alkyloxy(-CH₂-CH₂-O)_n-.
- 20 Preferred compounds are:
 - (±)-N-[1-(2-aminoethyl)-4-piperidinyl]-4-methyl-1-[1-(8-quinolinyl)ethyl]-IH-benzimidazol-2-amine monohydrate;
 - (\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(2-bromo-5,6,7,8-tetrahydro-8-quinolinyl)-IH-benzimidazol-2-amine trihydrochloride trihydrate;
- 25 (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-[(2-ethoxyethoxy)-8-quinolinyl-methyl]-4-methyl-1H-benzimidazol-2-amine;
 - (\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(2-chloro-5,6,7,8-tetrahydro-5-quinoxalinyl)-1H-benzimidazol-2-amine trihydrochloride trihydrate;
 - $(\pm)-N-[1-(2-a\min o-3-methylb\mu tyl)-4-piperidinyl]-1-[(1-methyl-1H-benzimidazol-4-methylbutyl)-4-piperidinyl]-1-[(1-methylbutyl)-1-[(1-methylbutyl)-1-[(1-methylbutyl)-1-[(1-methylbutyl)-1-[(1-methylbutyl)-1-[(1-methylbutyl)-1-[(1-methylbutyl)-1-[(1-methylbutyl)-1-[(1-methylbutyl)-1-[(1-methylbutyl)-1-[(1-methylb$
- 30 yl)methyl]-1H-benzimidazol-2-amine;
 - (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(ethoxy-8-quinolinylmethyl)-1H-benzimidazol-2-amine;
 - (\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-4-methyl-1-(5,6,7,8-tetrahydro-5-quinoxalinyl)-1H-benzimidazol-2-amine;
- 35 (\pm)-N-[1-(2-aminoethyl)-4-piperidinyl]-7-methyl-3-(8-quinolinylmethyl)-3H-imidazo-[4,5-b]pyridin-2-amine tetrahydrochloride trihydrate;
 - (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-7-methyl-3-(8-quinolinylmethyl)-3H-imidazo[4,5-b]pyridin-2-amine tetrahydrochloride monohydrate;

2-amine;

- (\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(8-quinolinylmethyl)-1H-imidazo[4,5-c]pyridin-2-amine trihydrochloride dihydrate; N-[1-(2-aminoethyl)-4-piperidinyl]-4-methyl-1-(8-quinolinylmethyl)-1H-benzimidazol-
- 5 *N*-[1-(8-quinolinylmethyl)-*1H*-benzimidazol-2-yl]-1,3-propanediamine trihydrochloride monohydrate:
 - (\pm)-N-[1-(2-aminoethyl)-4-piperidinyl]-1-[(2-ethoxyethoxy)-8-quinolinylmethyl]-IH-benzimidazol-2-amine trihydrochloride dihydrate;
 - $(\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(8-quinolinylmethyl)-1H-imidazo-1-(8-quinolinylmethyl)-1H-imidazo-1-(8-quinolinylmethyl)-1H-imidazo-1-(8-quinolinylmethyl)-1-(8-quinolinylmethyl)-1H-imidazo-1-(8-quinolinylmethyl)-1-(8-quinolinylmethylmethyl)-1-(8-quinolinylmethylmet$
- 10 [4,5-b]pyridine-2-amine trihydrochloride dihydrate;
 - (\pm)-N-[1-[1-(aminomethyl)-2-methylpropyl]-4-piperidinyl]-1-[(2-ethoxyethoxy)-8-quinolinylmethyl]-1H-benzimidazol-2-amine;
 - (±)-*N*-[1-(2-aminoethyl)-4-piperidinyl]-3-(2-quinolinylmethyl)-3*H*-imidazo-[4,5-b]pyridin-2-amine trihydrochloride trihydrate;
- (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(1-isoquinolinylmethyl)-1H-benzimidazol-2-amine trihydrochloride trihydrate;
 N-[1-(2-aminoethyl)-4-piperidinyl]-1-(5,6,7,8-tetrahydro-5-quinoxalinyl)-1H-benzimidazol-2-amine trihydrochloride trihydrate;
 the prodrugs, the N-oxides, the addition salts, the quaternary amines, the metal
 complexes and the stereochemically isomeric forms thereof.
 - Most preferred compounds are:
 - (\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-3-(quinolinylmethyl)-3H-imidazo[4,5-b]pyridin-2-amine;
- (\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-4-methyl-1-(8-quinolinylmethyl)-1H-benzimidazol-2-amine;
 - (\pm)-N-[1-(2-aminoethyl)-4-piperidinyl]-1-(2-chloro-5,6,7,8-tetrahydro-5-quinoxalinyl)-4-methyl-IH-benzimidazol-2-amine trihydrochloride.trihydrate;
 - (\pm) -N-[1-(2-aminoethyl)-4-piperidinyl]-1-(5,6,7,8-tetrahydro-2,3-dimethyl-5-
- quinoxalinyl)-1H-benzimidazol-2-amine trihydrochloride trihydrate; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-[(2-ethoxyethoxy)-8-quinolinyl-methyl]-1H-benzimidazol-2-amine;
 - (\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl-1-(3-chloro-5,6,7,8-tetrahydro-5-quinoxalinyl)-IH-benzimidazol-2-amine trihydrochloride monohydrate;
- 35 (±)-N-[1-(2-aminoethyl)-4-piperidinyl]-1-(3-chloro-5,6,7,8-tetrahydro-5-quinoxalinyl)-4-methyl-1H-benzimidazol-2-amine trihydrochloride dihydrate;

(\pm)-N-[1-(2-aminoethyl)-4-piperidinyl]-1-[(2-ethoxyethoxy)-8-quinolinylmethyl]-4-methyl-IH-benzimidazol-2-amine monohydrate;

(\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-3-(8-quinolinylmethyl)-3*H*-imidazo-[4,5-c]pyridin-2-amine trihydrochloride tetrahydrate;

5 (±)-N-[1-(2-aminoethyl)-4-piperidinyl]-3-(8-quinolinylmethyl)-3H-imidazo[4,5-b]-pyridin-2-amine;

(\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-4-methyl-1-[(1-methyl-1H-benzimidazol-4-yl)methyl]-1H-benzimidazol-2-amine;

(\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(2-chloro-5,6,7,8-tetrahydro-5-quinoxalinyl)-4-methyl-1H-benzimidazol-2-amine;

the prodrugs, the N-oxides, the addition salts, the quaternary amines, the metal complexes and the stereochemically isomeric forms thereof.

In general, compounds of formula (I) can be prepared by reacting an intermediate of formula (II-a) or (II-b), wherein P represents a protecting group, such as, for example C₁₋₄alkyloxycarbonyl, or those protecting groups mentioned in Chapter 7 of 'Protective Groups in Organic Synthesis' by T Greene and P. Wuyts (John Wiley & Sons Inc., 1991), with an intermediate of formula (III), wherein W₁ is a suitable leaving group, such as a halo atom, e.g. chloro, bromo, in the presence of a suitable base, such as, e.g. sodium hydride. Said reaction can be performed in a reaction-inert solvent, such as N,N-dimethylformamide.

$$Q = \begin{pmatrix} \mathbf{R}^{1} & \mathbf{G} - \mathbf{W}_{1} \\ \mathbf{Q} & \mathbf{A}^{1} & \mathbf{A}^{2} \\ \mathbf{Q} & \mathbf{A}^{2} & \mathbf{A}^{2} & \mathbf{A}^{2} \\ \mathbf{Q} & \mathbf{A}^{2} & \mathbf{A}^{2} & \mathbf{A}^{2} \\ \mathbf{Q} & \mathbf{A}^{2} & \mathbf{A}^{2} \\ \mathbf{Q} & \mathbf{A}^{2} & \mathbf{A}^{2} & \mathbf{A}^{2} & \mathbf{A}^{2} \\ \mathbf{Q} & \mathbf{A}^{2} & \mathbf{A}^{2} & \mathbf{A}^{2} \\ \mathbf{Q} & \mathbf{A}^{2} & \mathbf{A}^{2} &$$

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Compounds of formula (I) wherein, in the definition of Q, R² or at least one R⁶ substituent is hydrogen, said Q being represented by H-Q₁, and said compounds being represented by formula (I-a), can be prepared by deprotecting an intermediate of formula (IV) wherein P represents a protecting group, for example C₁.

4alkyloxycarbonyl, benzyl, or those protecting groups mentioned in Chapter 7 of

'Protective Groups in Organic Synthesis' by T Greene and P. Wuyts (John Wiley & Sons Inc., 1991).

$$P = Q_{1} = \begin{bmatrix} R^{1} & & & & \\ & & &$$

When P represents, for example, C1-4alkyloxycarbonyl, said deprotection reaction can be performed by, for example, acidic hydrolysis in the presence of a suitable acid, such 5 as hydrobromic, hydrochloric, sulfuric, acetic, or trifluoroacetic acid or a mixture of said acids, or by alkaline hydrolysis in the presence of a suitable base, such as, for example potassium hydroxide, in a suitable solvent such as water, alcohol, a mixture of water-alcohol, methylene chloride. Suitable alcohols are methanol, ethanol, 2-propanol, 1-butanol and the like. In order to enhance the rate of the reaction, it is 10 advantageous to heat the reaction mixture, in particular up to the reflux temperature. Alternatively, when P represents, for example, benzyl, the deprotection reaction can be performed by catalytic hydrogenation in the presence of hydrogen and an appropriate catalyst in a reaction-inert solvent. A suitable catalyst in the above reaction is, for example, platinum-on-charcoal, palladium-on-charcoal, and the like. An appropriate 15 reaction-inert solvent for said reaction is, for example, an alcohol, e.g. methanol, ethanol, 2-propanol and the like, an ester, e.g. ethylacetate and the like, an acid, e.g. acetic acid and the like.

The catalytic hydrogenation reaction described above can also be used to prepare a compound of formula (I-a) by deprotecting and reducing an intermediate of formula (IV) wherein Q₁ comprises an unsaturated bond, said Q₁ being represented by Q_{1a}(CH=CH), and said intermediate being represented by formula (IV-a).

Compounds of formula (I) wherein, in the definition of Q, both R⁶ substituents are hydrogen or R² and R⁴ are both hydrogen, said Q being represented by H₂N-Q₂, and said compounds being represented by formula (I-a-1), can also be prepared by deprotecting an intermediate of formula (V).

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$$Q_{2} = \begin{pmatrix} R^{1} & & & \\ & Q_{2} & & \\ & & &$$

Said deprotection reaction can be performed in the presence of a suitable base such as, for example hydrazine, or in the presence of a suitable acid, such as hydrochloric acid and the like, in a suitable solvent, such as an alcohol, acetic acid and the like.

Compounds of formula (I-a-1) can also be prepared by deprotecting an intermediate of formula (VI) according to the procedure described for the preparation of compounds of formula (I-a).

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(VIII)

Compounds of formula (I-a) or (I-a-1), wherein Q₁ or Q₂ comprise a hydroxy substituent, said Q₁ or Q₂ being represented by Q₁·(OH) or Q₂·(OH), and said compounds being represented by formula (I-a-2) or (I-a-1-1), can be prepared by deprotecting an intermediate of formula (VII) or (VIII) as described hereinabove for the preparation of compounds of formula (I-a).

(I-a-1-1)

$$P = Q_{1} \cdot (OP) \longrightarrow \begin{pmatrix} R^{1} & & & \\ & A^{2} &$$

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Compounds of formula (I) wherein, in the definition of Q, both R⁶ substituents are hydrogen or R² and R⁴ are both hydrogen, and the carbon adjacent to the nitrogen carrying the R⁶, or R² and R⁴ substituents contains at least one hydrogen, said Q being represented by H₂N-Q₃H, and said compounds being represented by formula (I-a-1-2) can also be obtained by reductive amination of intermediates of formula (IX) in the presence of a suitable amination reagent, such as, for example, ammonia, hydroxylamine, or benzylamine, and in the presence of a suitable reducing agent, e.g. hydrogen, and an appropriate catalyst. An appropriate catalyst in the above reaction is, for example, platinum-on-charcoal, palladium-on-charcoal, rhodium-on-Al₂O₃, and the like, optionally in the presence of a catalyst poison, such as a thiophene solution. A suitable reaction-inert solvent for the above reaction is, for example, an alcohol, e.g. methanol, ethanol, 2-propanol and the like.

$$(O\Longrightarrow)Q_3 \xrightarrow{R^1} a_1 \xrightarrow{a^1 = a^2} amination$$

$$(IX) \qquad H_2N - Q_3H \xrightarrow{N \qquad a^1 = a^2} a^3$$

Compounds of formula (I), wherein Q comprises a -CH₂NH₂ moiety, said Q being represented by H₂N-CH₂-Q₄, and said compounds being represented by formula (I-a-1-3) can be prepared by reducing an intermediate of formula (X).

NC-Q₄

$$\stackrel{\text{R}^1}{\underset{\text{a}}{\bigvee}}$$
 $\stackrel{\text{reduction}}{\underset{\text{a}}{\bigvee}}$
 $\stackrel{\text{reduction}}{\underset{\text{a}}{\bigvee}}$
 $\stackrel{\text{R}^1}{\underset{\text{a}}{\bigvee}}$
 $\stackrel{\text{a}^1}{\underset{\text{a}}{\bigvee}}$
 $\stackrel{\text{a}^1}{\underset{\text{a}}{\bigvee}}$

Said reduction can be performed with a suitable reducing agent, such as lithium aluminium hydride or hydrogen, optionally in the presence of a suitable catalyst, such as Raney Nickel. A suitable solvent for the above reaction is, for example, tetrahydrofuran, or a solution of ammonia in an alcohol. Suitable alcohols are methanol, ethanol, 2-propanol and the like. Said reduction reaction performed in a solution of ammonia in an alcohol can also be used to prepare compounds of formula (I-a-1-3), wherein R^1 is substituted with C_{1-6} alkyloxy C_{1-6} alkyl, said R^1 being represented by $R^{1'}$ - C_{1-6} alkyloxy C_{1-6} alkyl, and said compounds being represented by formula (I-a-1-3-1) starting from an intermediate of formula (X-a).

Compounds of formula (I), wherein Q comprises a -CH₂-CHOH-CH₂-NH₂ moiety, said Q being represented by H₂N-CH₂-CHOH-CH₂-Q₄, and said compounds being represented by formula (I-a-1-3-2), can be prepared by reacting an intermediate of formula (XI) with ammonia in the presence of a suitable reaction-inert solvent, such as an alcohol, e.g. methanol.

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Compounds of formula (I), wherein, in the definition of Q, R² or one R⁶ substituent is formyl, said Q being represented by H-C(=O)-Q₁, and said compounds being represented by formula (I-b), can be prepared by reacting an intermediate of formula (XII) with formic acid, formamide and ammonia.

$$C_{1^{-4}}\text{alkyl} - C_{1^{-4}}\text{alkyl} - C_{1^{-$$

Compounds of formula (I), wherein, in the definition of Q, R² is other than hydrogen, said R² being represented by R^{2a}, R⁴ is hydrogen, and the carbon atom adjacent to the nitrogen atom carrying the R² and R⁴ substituents, carries also at least one hydrogen atom, said Q being represented by R^{2a}-NH-HQ₅, and said compounds being represented by formula (I-c), can be prepared by reductive amination of an intermediate of formula (XIII) with an intermediate of formula (XIV) in the presence of a suitable reducing agent, such as hydrogen, and a suitable catalyst, such as palladium-on-charcoal, platinum-on-charcoal, and the like. A suitable reaction-inert solvent for the above reaction is, for example, an alcohol, e.g. methanol, ethanol, 2-propanol and the like.

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$$(O=)Q_{5} \xrightarrow{R^{1}} A_{3}^{2} + R^{2a} \xrightarrow{NH_{2}} A_{3}^{2} + R^{2a} \xrightarrow{NH_{2}} A_{3}^{2}$$

$$(XIII) \qquad (XIV)$$

Compounds of formula (I-c), wherein R^{2a} represents C_{1-10} alkyl substituted with $N(R^6)_2$ and with hydroxy, and the carbon atom carrying the hydroxy, carries also two hydrogen atoms, said R^{2a} being represented by $[(C_{1-9}alkyl)CH_2OH]-N(R^6)_2$, and said compounds being represented by formula (I-c-1), can be prepared by reducing an intermediate of formula (XV) in the presence of a suitable reducing agent, such as lithium aluminium hydride, in a suitable reaction-inert solvent, such as tetrahydrofuran.

$$(R^{6})_{2}N-(C_{1}-9alkyl)-NH-HQ_{5}-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}N-(R^{6})_{2}$$

Compounds of formula (I) wherein, in the definition of Q, R² or one R⁶ substituent is hydrogen, said Q being represented by H-Q₁, and wherein R¹ is a bicyclic heterocycle substituted with 1 or more substituents selected from hydroxy, hydroxyC₁₋₆alkyl, or HO(-CH₂-CH₂-O)_n-, said substituents being represented by formula A-OH, said R¹ being represented by R^{1a}-(A-OH)_w, with w being the amount of substituents on R^{1a} ranging from 1 to 4, and said compounds being represented by formula (I-d), can be prepared by deprotecting an intermediate of formula (XVI) with a suitable acid, such as hydrochloric acid and the like, optionally in the presence of a suitable solvent, such as an alcohol. Suitable alcohols are methanol, ethanol, 2-propanol and the like. Alternatively, one protecting group may also protect more than one substituent of R^{1a}, said protecting group being represented by P₁, as represented by formula (XVI-a). The two ways of protecting the substituents of R^{1a}, i.e. with a separate, as in formula (XVI), or a combined, as in formula (XVI-a), protecting group, may also be combined in the same intermediate, as represented by formula (XVI-b).

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Compounds of formula (I), wherein Q is a radical of formula (b-2), said compounds

being represented by formula (I-e), can be prepared by reacting an intermediate of
formula (XVII) with an intermediate of formula (XVIII) in the presence of sodium
cyanide and a suitable reaction-inert solvent, such as an alcohol, e.g. methanol and the
like.

$$C_{1^{-4}alkyl} - O - C_{-Alk} - X^{1} - Alk - X^{1} - A$$

Compounds of formula (I), wherein in the definition of Q, X² is C₂₋₄alkyl-NR⁴, said Q being represented by Q₆N-CH₂-C₁₋₃alkyl-NR⁴, and said compounds being represented by formula (I-p), can be prepared by reacting an intermediate of formula (XIX) with an intermediate of formula (XX) in the presence of isopropyl titanate (IV) and a suitable

reducing agent, such as NaBH₃CN, and in the presence of a suitable reaction-inert solvent, such as methylene chloride and an alcohol, e.g. ethanol.

$$H = C - C_{1-3} \text{alkyl} - NR^4$$

$$N = A^{1} A^{2} + Q_6 N - CH_2 - C_{1-3} \text{alkyl} - NR^4$$

$$N = A^{1} A^{2} A^{2} + Q_6 N - CH_2 - C_{1-3} \text{alkyl} - NR^4$$

$$N = A^{1} A^{2} A^{2} A^{3}$$

$$(XIX) \qquad (XX) \qquad (i-p)$$

Compounds of formula (I-p), wherein R^2 is C_{1-6} alkylcarbonyl, and Q is a radical of formula (b-6), wherein Y^1 is NR^2 , said compounds being represented by formula (I-p-1), can be prepared by reacting an intermediate of formula (XIX) with an intermediate of formula (XX-a) according to the procedure described for the preparation of a compound of formula (I-p).

(I-p-1)

Compounds of formula (I), wherein G is substituted with hydroxy or HO(-CH₂CH₂O)_n-, said G being represented by G₁-OH, and said compounds being represented by formula (I-q), may be prepared by deprotecting an intermediate of formula (XXI), wherein P represents a suitable protecting group, for example, benzyl. Said deprotection reaction can be performed by catalytic hydrogenation in the presence of hydrogen and an
 appropriate catalyst in a reaction-inert solvent. A suitable catalyst in the above reaction is, for example, platinum-on-charcoal, palladium-on-charcoal, and the like. An appropriate reaction-inert solvent for said reaction is, for example, an alcohol, e.g. methanol, ethanol, 2-propanol and the like, an ester, e.g. ethylacetate and the like, an acid, e.g. acetic acid and the like.

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$$P \longrightarrow G_1$$

$$Q \longrightarrow N$$

$$A = A^2$$

$$A = A^3$$

$$A = A^$$

Compounds of formula (I), wherein G is substituted with hydroxy and the carbon atom carrying the hydroxy substituent carries also at least one hydrogen, said G being represented by H-G₂-OH, and said compounds being represented by formula (I-q-1), can also be prepared by reducing an intermediate of formula (XXII).

$$(O=)G_2$$

$$O=(O=)G_2$$

$$O=(O=)G_3$$

$$O=(O=)G_4$$

$$O=(O=)G_4$$

$$O=(O=)G_4$$

$$O=(O=)G_4$$

$$O=(O=)G_4$$

$$O=(O=)G_4$$

$$O=(O=)$$

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Said reduction reaction can be performed in the presence of a suitable reducing agent, such as, for example sodium borohydride, in a reaction-inert solvent, such as an alcohol or tetrahydrofuran or a mixture thereof. Suitable alcohols are methanol, ethanol, 2-propanol and the like.

Compounds of formula (I) may be converted into each other following art-known functional group transformation reactions, comprising those described hereinafter.

The compounds of formula (I) may be converted to the corresponding N-oxide forms following art-known procedures for converting a trivalent nitrogen into its N-oxide form. Said N-oxidation reaction may generally be carried out by reacting the starting material of formula (I) with an appropriate organic or inorganic peroxide. Appropriate inorganic peroxides comprise; for example, hydrogen peroxide, alkali metal or earth alkaline metal peroxides, e.g. sodium peroxide, potassium peroxide; appropriate organic peroxides may comprise peroxy acids such as, for example, benzenecarboperoxoic acid or halo substituted benzenecarboperoxoic acid, e.g. 3-chlorobenzenecarboperoxoic acid, peroxoalkanoic acids, e.g. peroxoacetic acid, alkylhydroperoxides, e.g. t.butyl hydro-peroxide. Suitable solvents are, for example, water, lower alcohols, e.g. ethanol and the like, hydrocarbons, e.g. toluene, ketones, e.g. 2-butanone, halogenated hydrocarbons, e.g. dichloromethane, and mixtures of such solvents.

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Compounds of formula (I), wherein R^1 is a bicyclic heterocycle substituted with $C_{1\text{-}6}$ alkyloxycarbonyl, said R^1 being represented by R^1 - $C(=0)OC_{1\text{-}6}$ alkyl, and said compounds being represented by formula (I-f), can be prepared by esterification of a compound of formula (I-g) in the presence of a suitable alcohol, e.g. methanol, ethanol, propanol, butanol, pentanol, hexanol and the like, and in the presence of a suitable acid, such as hydrochloric acid and the like.

Q
$$= \frac{R^{1}-C(=0)OH}{a^{1}-a^{2}}$$
 esterification $= \frac{R^{1}-C(=0)OC_{1-6}alkyl}{N}$ $= \frac{a^{1}-a^{2}}{N}$ $= \frac{a^{1}-a^{2}}{a^{3}}$ $= \frac{a^{1}-a^{2}}{N}$ $= \frac{a^{1}-a^{2}}{N$

Compounds of formula (I-a) may be converted into compounds of formula (I) wherein, in the definition of Q, R^2 or at least one R^6 substituent is other than hydrogen, said R^2 or R^6 being represented by Z_1 , said Q being represented by Z_1 - Q_1 , and said compounds being represented by formula (I-h), by reaction with a reagent of formula (XXIII), wherein W_2 is a suitable leaving group, such as a halo atom, e.g. bromo, or 4-methylbenzenesulphonate, in the presence of a suitable base, such as, for example disodium carbonate, dipotassium carbonate, sodium hydroxide and the like, in a reaction-inert solvent, e.g. 3-methyl-2-butanone, acetonitrile, N,N-dimethylformamide.

Compounds of formula (I-h), wherein, in the definition of Z_1 , R^2 is CH_2 - $C_{1.9}$ alkyl substituted with $N(R^6)_2$, said compounds being represented by formula (I-h-1), can also be prepared by reacting a compound of formula (I-a) wherein, in the definition of H-Q₁, R^2 is hydrogen, said H-Q₁ being represented by H-Q_{1b}, and said compounds being represented by formula (I-a-3), with an intermediate of formula (XXIV), in the presence of a suitable reducing agent, such as sodium cyanoborohydride, in a suitable reaction-inert solvent, such as an alcohol.

Compounds of formula (I-h), wherein Z₁ comprises formyl, C₁₋₆alkylcarbonyl, Hetcarbonyl or C₁₋₆alkyloxycarbonyl, said Z₁ being represented by Z_{1a}, and said compounds being represented by formula (I-h-2), can be converted into compounds of formula (I-a), by acidic hydrolysis in the presence of a suitable acid, such as hydrobromic, hydrochloric, sulfuric, acetic, or trifluoroacetic acid or a mixture of said acids, or by alkaline hydrolysis in the presence of a suitable base, such as, for example potassium hydroxide, in a suitable solvent such as water, alcohol, a mixture of water-alcohol, methylene chloride. Suitable alcohols are methanol, ethanol, 2-propanol, 1-butanol, sec. butanol and the like. In order to enhance the rate of the reaction, it is advantageous to work at elevated temperatures.

$$Z_{1a} = Q_{1} + Q_{$$

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Compounds of formula (I-b) can be prepared by reacting a compound of formula (I-a) with formic acid.

Compounds of formula (I) wherein R^1 is a bicyclic heterocycle substituted with hydroxy, said R^1 being represented by HO- R^1 , and said compounds being represented by formula (I-i), can be prepared by deprotecting a compound of formula (I-j), wherein R^1 is a bicyclic heterocycle substituted with C_{1-6} alkyloxy or aryl C_{1-6} alkyloxy, said C_{1-6} alkyl or aryl C_{1-6} alkyl being represented by Z_2 , and said R^1 being represented by Z_2 -O- R^1 . Said deprotection can be performed in a reaction-inert solvent, such as, for

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example methylene chloride, in the presence of a suitable deprotecting agent, e.g. tribromoborane.

Compounds of formula (I) wherein R¹ is a bicyclic heterocycle substituted with halo(-CH₂-CH₂-O)_n, said compounds being represented by formula (I-k), can be converted into compounds of formula (I-l-1) or (I-l-2) by reaction with an appropriate amine of formula (XXV) or (XXVI) in a suitable reaction-inert solvent, e.g. tetrahydrofuran.

$$(C_{1-6}alkyl)N (-CH_{2}-CH_{2}-O)_{n} = R^{1}$$

$$(C_{1-6}alkyl)N (-CH_{2}-CH_{2}-O)_{n} = R^{1}$$

$$(XXV)$$

$$(I-l-1)$$

$$(C_{1-6}alkyl)_{2}N (-CH_{2}-CH_{2}-O)_{n} = R^{1}$$

$$(C_{1-6}alkyl)_{2}N (-CH_{2}-CH_{2}-O)_{n} = R^{1}$$

$$(XXV)$$

$$(I-l-1)$$

$$(C_{1-6}alkyl)_{2}N (-CH_{2}-CH_{2}-O)_{n} = R^{1}$$

$$(XXVI)$$

$$(I-l-2)$$

Compounds of formula (I) wherein R¹ is a bicyclic heterocycle substituted with halo, said compounds being represented by formula (I-m) can be converted into compounds of formula (I) by reaction with 1-butanethiol in the presence of palladium-on-charcoal and CaO in a suitable reaction-inert solvent, such as tetrahydrofuran.

Q
$$R^{1'}$$
 Q $R^{1'}$ Q R^{1} R^{1} R^{1} R^{1} R^{1} R^{2} R^{1} R^{2} R^{1} R^{2} R^{2

Compounds of formula (I) wherein a hydrogen atom in the radicals of formula (a-1), (a-2), (a-3), (a-4) or (a-5) is replaced by nitro, said compounds being represented by formula (I-n) may be reduced to a compound of formula (I-o) in the presence of a suitable reducing agent, such as hydrogen, optionally in the presence of a suitable catalyst, such as platinum-on-charcoal, and optionally in the presence of a suitable catalyst poison, e.g. a thiophene solution. The reaction may be performed in a suitable reaction-inert solvent, such as an alcohol.

In the following paragraphs, there are described several methods of preparing the intermediates in the foregoing preparations. A number of intermediates and starting materials are commercially available or are known compounds which may be prepared according to conventional reaction procedures generally known in the art or analogous to the procedures described in EP-A-0005318, EP-A-0099139, EP-A-0151824, EP-A-0151826, EP-A-0232937, EP-A-0295742, EP-A-0297661, EP-A-0539420, EP-A-0539421, US 4,634,704, US 4,695,569.

In the foregoing and the following preparations, the reaction mixture is worked up following art-known methods and the reaction product is isolated and, if necessary, further purified.

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Intermediates of formula (III) can be prepared by reacting an intermediate of formula (XXVII) with a suitable leaving group, i.e. W₁, introducing agent, e.g. 1-halo-2,5-pyrrolidinedione in the presence of dibenzoyl peroxide, in a reaction-inert solvent, e.g. tetrachloromethane.

$$R^{1}$$
— G — H
 O
 R^{1} — G — W_{1}

(XXVII)

 (III)

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Intermediates of formula (XXVII), wherein R¹ is a bicyclic heterocycle substituted with chloro, said R¹ being represented by Cl-R¹ and said intermediates being represented by formula (XXVII-a) can be prepared by reacting an intermediate of formula (XXVIII),

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wherein (O=)R^{1b}H is defined as a carbonyl derivative of R^{1'} wherein one carbon or nitrogen, adjacent to the carbonyl, carries at least one hydrogen, with phosphorus oxychloride. Intermediates of formula (XXVIII) may also react as their enol tautomeric forms.

$$(O=)R^{1b}H - G - H \qquad POCl_3$$

$$(XXVIII) \qquad (XXVIII-a)$$

Intermediates of formula (XXVII), wherein R¹ is 2-trifluoromethyl-3-methyl (3H)-imidazo[4,5-b]pyridine, and G is CH₂, said intermediates being represented by formula (XXVII-b), can be prepared by reacting N-2,6-dimethyl-2,3-pyridinediamine (Heterocycles, 38, p 529, 1994), with trifluoroacetic acid.

$$\begin{array}{c} H \\ CH_2 \\ NH_2 \end{array} \qquad \begin{array}{c} CF_3\text{-COOH} \\ \end{array} \qquad \begin{array}{c} H \\ CH_2 \\ N \end{array} \qquad \begin{array}{c} N \\ N \end{array} \qquad \begin{array}{c} CF_3 \\ \end{array}$$

Intermediates of formula (III) wherein W_1 is chloro, which is attached to a carbon atom carrying at least one hydrogen, said G being represented by G_3H , and said intermediates being represented by formula (III-a) can also be prepared by reacting an intermediate of formula (XXIX) with thionylchloride in a reaction-inert solvent, e.g.

15 methylenechloride.

$$R^1$$
— G_3H — OH $SOCI_2$ R^1 — G_3H — CI (XXIX) (III-a)

Intermediates of formula (XXIX) can be prepared by reducing an intermediate of formula (XXX) in a reaction-inert solvent, e.g. an alcohol, in the presence of a suitable reducing agent, e.g. sodium borohydride.

$$R^1$$
 $G_3(=O)$ R^1 G_3H OH (XXX)

Alternatively, intermediates of formula (XXIX) can also be prepared by deprotecting an intermediate of formula (XXXI), wherein P is a suitable protecting group, e.g. C₁₋₄alkylcarbonyl, in a reaction-inert solvent, such as an alcohol, in the presence of a suitable base, e.g. sodium hydroxide.

$$R^1$$
— G_3H — O — P R^1 — G_3H — OH (XXIX)

Intermediates of formula (XXX), wherein $G_3(=0)$ is CH(=0), said intermediates being represented by formula (XXX-a), can be prepared by reacting an intermediate of formula (XXXII), wherein W_3 is a suitable leaving group, such as a halo atom, e.g. bromo, with N_1N_2 -dimethylformamide in the presence of butyllithium in a reaction-inert solvent, e.g. tetrahydrofuran, diethylether or a mixture thereof.

$$R^{\perp}W_3$$
 $R^{\perp}CH(=0)$ (XXX-a)

Intermediates of formula (XXX-a) can also be prepared by oxidizing an intermediate of formula R¹-CH₂-OH in the presence of a suitable oxidizing agent, e.g. MnO₂ in a reaction-inert solvent, e.g. methylenechloride.

$$R^1$$
— CH_2 — OH R^1 — $CH(=O$ (XXX-a)

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Intermediates of formula R¹-CH₂-OH, wherein R¹ is 2,3-dimethylquinoxaline, said intermediates being represented by formula (XCI) can be prepared by reducing an intermediate of formula (XCII) in a reaction-inert solvent, e.g. tetrahydrofuran, in the presence of a suitable reducing agent, e.g. potassium borohydride in the presence of lithium chloride.

Intermediates of formula (XCII) can be prepared by reacting ethyl 2,3-diaminobenzoate (Tetrahydron, 28, 3271, 1972) with 2,3-butanedione in the presence of disodium disulfite.

$$\begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \\ \\ \end{array} \end{array} \\ \begin{array}{c} \\ \end{array} \end{array} \begin{array}{c} \\ \\ \end{array} \end{array} \begin{array}{c} \\ \\ \end{array} \begin{array}{c} \\ \\ \end{array} \begin{array}{c} \\ \end{array} \begin{array}{c} \\ \\ \end{array} \end{array} \begin{array}{c} \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\$$

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Intermediates of formula (XXXI), wherein R^1 is 5,6,7,8-tetrahydroquinoline, which can optionally be substituted, G_3H is CH_2 , and P is C_{1-4} alkylcarbonyl, said intermediates being represented by formula (XXXI-a) can be prepared by reacting an intermediate of formula (XCIII) with C_{1-4} alkylacid anhydride at elevated temperatures in the presence of a suitable base, e.g. sodium hydroxide.

$$\begin{array}{c|c} CH_3 & CH_2 & CCH_2 & CC_{1^-4}alkyl \\ \hline \\ CCH_2 & CC_{1^-4}alkyl \\ \hline \\ CCH_2 & CC_{1^-4}alkyl \\ \hline \\ CCC_{1^-4}alkyl \\ \hline \\$$

Intermediates of formula (XCIII) can be prepared by oxidizing an intermediate of formula (XCIV) with a suitable oxidizing agent, e.g. a peroxide such as 3-chlorobenzenecarboperoxoic acid, in a reaction-inert solvent, e.g. methylene chloride.

Intermediates of formula (XCIV) can be prepared by reducing an intermediate of formula (XCV) (Org. Prep. Proced. Int., 23, p 386-387, 1991) with an appropriate reducing agent, e.g. hydrogen, in the presence of a suitable catalyst, e.g. palladium-on-charcoal, and a suitable acid, e.g. trifluoroacetic acid.

Intermediates of formula (IV) can be prepared by reacting an intermediate of formula (XXXIII-a) or (XXXIII-b), wherein P represents a suitable protecting group, such as, for example, C₁₋₄alkyloxycarbonyl, with an intermediate of formula (III) according to the reaction described for the general preparation of compounds of formula (I).

Intermediates of formula (IV) can also be prepared by reacting an intermediate of formula (XXXIII-a) with an intermediate of formula (XXXIV) that has reacted with methanesulfonyl chloride, in the presence of a suitable base, such as sodium hydride, and in the presence of a suitable reaction-inert solvent, e.g. N,N-dimethylformamide.

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$$P = Q_1 = \begin{bmatrix} H & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & &$$

Intermediates of formula (IV) can also be prepared by a cyclization reaction of an intermediate of formula (XXXV) in a reaction-inert solvent, e.g. an alcohol or N,N-dimethylformamide, in the presence of mercury oxide and sulphur.

Intermediates of formula (IV) wherein Q_1 comprises an unsaturated bond, said Q_1 being represented by $Q_{1a}(CH=CH)$, and said intermediates by formula (IV-a), can be prepared by reacting an intermediate of formula (XXXVI) with an intermediate of formula (III) in the presence of a suitable base, such as dipotassium carbonate.

Intermediates of formula (IV) wherein, in the definition of Q_1 , the X^1 or X^2 moieties in the radicals of formula (b-1) to (b-8) represent NH, said Q_1 being represented by Q_{1c} -NH, and said intermediates by formula (IV-b), may also be prepared by reacting an intermediate of formula (XXXVII) with an intermediate of formula (XXXVIII).

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Intermediates of formula (IV) wherein R¹ is a bicyclic heterocycle substituted with amino or mono- or di(C₁₋₆alkyl)amino, said R¹ being represented by R^{5a}R^{5b}N-R¹, wherein R^{5a} and R^{5b} are defined as described above, and said intermediates being represented by formula (IV-c), can be prepared by reacting an intermediate of formula (XXXIX) with an appropriate amine, represented by formula (XL), in the presence of an appropriate catalyst, e.g. palladium, and (R)-(+)-2,2'-bis(diphenyl-phosphino)-1,1'-binaphtyl, in a suitable reaction-inert solvent, e.g. tetrahydrofuran.

halo—
$$R^{1'}$$
 R^{5a}
 R^{5b}
 R^{5a}
 R^{5a}
 R^{5b}
 R^{5a}
 R^{5a}
 R^{5b}
 R^{5a}
 R^{5a}
 R^{5b}
 R^{5a}
 R^{5a}

Intermediates of formula (IV) wherein R¹ is a bicyclic heterocycle substituted with C(=O)-NR^{5a}R^{5b}, wherein R^{5a} and R^{5b} are defined as described above, said R¹ being represented by R^{5a}R^{5b}N-C(=O)-R¹, and said intermediates being represented by formula (IV-d), can be prepared by reacting an intermediate of formula (XXXIX) with an appropriate amine, represented by formula (XL), under an atmosphere of carbon monoxide, in the presence of a suitable catalyst, e.g. palladium (II) acetate, and

1,3-bis(diphenylphosphino)propane, in a suitable reaction-inert solvent, e.g. tetrahydrofuran.

halo—
$$R^{1'}$$
 R^{5b}
 R^{5a}
 R^{5b}
 R^{5a}
 R^{5b}
 R^{5a}
 R^{5b}
 R^{5a}
 R^{5a}

Intermediates of formula (IV) wherein P-Q₁ comprises C_{1-10} alkyl or C_{3-7} cycloalkyl substituted with NR⁶-P, said C_{1-10} alkyl or C_{3-7} cycloalkyl being represented by Z₃, said P-Q₁ being represented by P-NR⁶-Z₃-Q_{1b}, and said intermediates being represented by formula (IV-e), can be prepared by reacting a compound of formula (I-a-3) with an intermediate of formula (XLI), wherein W₄ represents a suitable leaving group, such as p-toluenesulphonate. Said reaction can be performed in a reaction-inert solvent, e.g. acetonitrile, in the presence of a suitable base, e.g. dipotassium carbonate.

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Intermediates of formula (IV-e), wherein R⁶ is hydroxyC₁₋₆alkyl, said intermediates being represented by formula (IV-e-1), can be prepared by reacting an intermediate of formula (XLII) with an intermediate of formula (XLIII) in the presence of a suitable base, e.g. dipotassium carbonate, and a suitable solvent, e.g. acetonitrile.

$$Q = \begin{bmatrix} Q & Q & Q \\ Q & Q \\ Q & Q \end{bmatrix}$$

$$Q = \begin{bmatrix} Q & Q \\ Q & Q \\ Q & Q \end{bmatrix}$$

$$Q = \begin{bmatrix} Q & Q \\ Q & Q \\ Q & Q \end{bmatrix}$$

$$Q = \begin{bmatrix} Q & Q \\ Q & Q \\ Q & Q \end{bmatrix}$$

$$Q = \begin{bmatrix} Q & Q \\ Q & Q \\ Q & Q \end{bmatrix}$$

$$Q = \begin{bmatrix} Q & Q \\ Q & Q \\ Q & Q \end{bmatrix}$$

$$Q = \begin{bmatrix} Q & Q \\ Q & Q \\ Q & Q \end{bmatrix}$$

$$Q = \begin{bmatrix} Q & Q \\ Q & Q \\ Q & Q \\ Q & Q \end{bmatrix}$$

$$Q = \begin{bmatrix} Q & Q \\ Q$$

Intermediates of formula (XXXIII-a) or (XXXIII-b) can be prepared by protecting an intermediate of formula (XLIV) with a suitable protecting group, such as, for example, C₁₋₄alkyloxycarbonyl, in a reaction-inert solvent, such as methylene chloride or an alcohol, e.g. methanol, ethanol, 2-propanol and the like, in the presence of a suitable

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reagent, e.g. di C₁₋₄alkyl dicarbonate and optionally in the presence of a suitable base, e.g. sodium acetate.

Alternatively, intermediates of formula (XXXIII-a) or (XXXIII-b) can be converted into an intermediate of formula (XLIV) by reaction with a suitable acid, such as hydrochloric acid or hydrobromic acid and the like or mixtures thereof, in the presence of a suitable solvent, e.g. water.

Intermediates of formula (XXXIII-a) or (XXXIII-b), wherein in the definition of Q₁, the X¹ or X² moieties in the radicals of formula (b-1) to (b-8) represent NH, said Q₁ being represented by Q_{1c}-NH, and said intermediates by formula (XXXIII-a-1) or (XXXIII-b-1), can be prepared by reacting an intermediate of formula (XLV-a) or (XLV-b), wherein W₅ represents a suitable leaving group, such as for example a halo atom, e.g. chloro, with an intermediate of formula (XLVI).

$$W_{5} = \begin{pmatrix} A_{1} & A_{2} & A_{3} & A_{4} & A_{3} & A_{4} & A_{4} & A_{3} & A_{4} & A_$$

Intermediates of formula (XLV-a) or (XLV-b) can be prepared by reacting an intermediate of formula (XLVII-a) or (XLVII-b) with $H_2P(=O)(W_5)_3$ in the presence of a suitable acid, e.g. hydrochloric acid.

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$$O = \bigvee_{\substack{N \\ N \\ N}} \bigvee_{\substack{a^1 \\ a^2 \\ a^3}} \bigvee_{\substack{a^2 \\ a^3 \\ a^3}} \bigvee_{\substack{a^1 \\ a^2 \\ a^3}} \bigvee_{\substack{a^1 \\ a^3 \\ a^3}}$$

Intermediates of formula (XLVII-a) or (XLVII-b) can be prepared by reacting an intermediate of formula (XLVIII-a) or (XLVIII-b) with an intermediate of formula (IL).

Intermediates of formula (XXXIII-a) can also be prepared by reacting an intermediate of formula (XLVIII-a) with P-Q₁-C(=NH)-O-CH₂-CH₃ in a reaction-inert solvent, such as an alcohol.

Intermediates of formula (XXXV) can be prepared by reacting an intermediate of formula (L) with an intermediate of formula P-Q₁=C=S, which is synthesized according to the procedures described in EP 0005318, in a reaction-inert solvent, such as an alcohol, e.g. ethanol. To increase the reaction rate, the reaction may be performed at elevated temperatures.

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$$\begin{array}{c} G - R^{1} \\ \downarrow \\ NH_{2} \\ \downarrow \\ NH_{2} \\ \downarrow \\ a^{3} \end{array} + P-Q_{1} = C = S \\ \begin{array}{c} P - Q_{1} - C - HN - \frac{1}{3} \\ \downarrow \\ S \\ \end{array}$$

$$(XXXV)$$

Intermediates of formula (L) can be obtained by reducing an intermediate of formula (LI) in a reaction-inert solvent, e.g. an alcohol, in the presence of a suitable reducing agent, e.g. hydrogen, and an appropriate catalyst, e.g. Raney Nickel.

Intermediates of formula (LI) can be prepared by reacting an intermediate of formula (LII) with an intermediate of formula (LIII), in which W₆ represents a suitable leaving group, such as a halo atom, e.g. chloro. This reaction may be performed in a reaction-inert solvent, e.g. acetonitrile, in the presence of a suitable base, e.g. dipotassium carbonate.

$$R^{1}$$
— G — NH_{2} + W_{6} A_{4} A_{3} A_{4} A_{3} A_{4} A_{3} A_{4} A_{3} A_{4} A_{3} A_{4} A_{3} A_{4} A_{4}

Intermediates of formula (LII) can be prepared by reacting an intermediate of formula (LIV) with a suitable acid, such as hydrochloric acid, in the presence of a suitable solvent, e.g. an alcohol, e.g. ethanol.

Intermediates of formula (LIV) can be prepared by reacting an intermediate of formula (III) with NaN[C(=0)H]₂.

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$$R^{1}-G-W_{1}$$
 + NaN[C(=O)H]₂ $R^{1}-G-N$ $C=O$
(III)

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Intermediates of formula (LI) can also be prepared by reacting an intermediate of formula (LIII) with an intermediate of formula (LV) (J. Org. Chem., 25, p 1138, 1960) in a reaction-inert solvent, e.g. N,N-dimethylformamide, in the presence of an appropriate base, e.g. sodium hydride.

$$R^{1}-G-NH-C-H + W_{0} = \begin{pmatrix} W_{0} & A_{1} & A_{2} & A_{3} & A_{4} & A_{3} & A_{4} & A_{3} & A_{4} & A$$

Intermediates of formula (XXXVI) can be prepared by dehydrating an intermediate of formula (LVI) with a suitable acid, such as sulfuric acid.

$$P = Q_{1a}(CH_2-CHOH) \longrightarrow N \longrightarrow A^{1 - 1} A^{2 -$$

Intermediates of formula (LVI) wherein, in the definition of Q_{1a}, the X¹ or X² moieties are CH₂, said Q_{1a} being represented by Q_{1a}, and said intermediates being represented by formula (LVI-a), can be prepared by reacting a carbonyl moiety of formula (LVII) with an intermediate of formula (LVIII) in the presence of N,N-disopropylamine and butyl lithium, in a suitable reaction-inert solvent, e.g. tetrahydrofuran.

$$P = Q_{1a}(CH_2-C=0) + CH_3 = Q_{1a}(CH_2-CHOH) = CH_2 = Q_{1a}(CH_2-CHOH) = CH_2 = Q_{1a}(CH_2-CHOH) =$$

Intermediates of formula (IV), wherein G is C_{1-10} alkanediyl substituted with C_{1-6} alkyloxy, aryl C_{1-6} alkyloxy, $HO(-CH_2CH_2O)_n$ -, C_{1-6} alkyloxy($-CH_2CH_2O)_n$ -, or aryl C_{1-6} alkyloxy($-CH_2CH_2O)_n$ -, said group of substituents being represented by $O-Z_4$, said G being represented by Z_4 - $O-G_1$, and said intermediates being represented by formula (IV-f), can be prepared by reacting an intermediate of formula (XXXIII-a), with an intermediate of formula (LIX), optionally in the presence of a suitable acid, such as p-toluenesulfonic acid and the like, and optionally in the presence of a suitable solvent,

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such as N,N-dimethylacetamide. To increase the reaction rate, the reaction may be carried out at elevated temperatures.

$$P = Q_1 = \begin{pmatrix} R^1 \\ Z_4 = O - Q_1 \\ N = A^2 = A^3 \end{pmatrix}$$

$$(XXXIII-a)$$

$$Z_4 = O - Q_1 \\ O - Z_4$$

$$P = Q_1 = \begin{pmatrix} R^1 \\ N = A^2 = A^3 \\ N = A^2 = A^3 \end{pmatrix}$$

$$(IV-f)$$

Intermediates of formula (LIX) can be prepared by reacting an intermediate of formula (LX) with a reagent of formula (LXI) or (LXII) in a reaction-inert solvent, such as an alcohol, or toluene, in the presence of an acid, e.g. 4-methylbenzenesulphonic acid.

$$Z_4$$
-O-H (LXI) or Z_4 -O-Z₄ (LX) Q -Z₄ Q -Q-C₁-4alkyl (LXII)

Intermediates of formula (LX) can be prepared by oxidizing an intermediate of formula (LXIII) with a suitable oxidizing agent, e.g. MnO₂, in a reaction-inert solvent, such as methylene chloride.

$$R^1$$
— G_1 H—OH R^1 — G_1 (=O) (LX)

Intermediates of formula (IV-f) can also be prepared by reacting an intermediate of formula (IV) wherein G is C_{1-10} alkanediyl substituted with hydroxy, said G being represented by G_1 -OH, and said intermediates being represented by formula (IV-g), with an intermediate of formula (LXIV), wherein W_7 is a suitable leaving group, such as a halo atom, e.g. iodo, in the presence of a suitable base, e.g. sodium hydride, in a reaction-inert solvent, e.g. tetrahydrofuran.

$$P = Q_1$$

$$|V - g|$$

Intermediates of formula (IV-g), wherein the carbon atom of G_1 carrying the hydroxy, also carries a hydrogen atom, said G_1 -OH being represented by H-G₂-OH, and said intermediates being represented by formula (IV-g-1), can be prepared by reducing an

intermediate of formula (LXV) in the presence of a suitable reducing agent, e.g. sodium borohydride, in a reaction-inert solvent, such as an alcohol, tetrahydrofuran or a mixture thereof. Intermediates of formula (LXV) can also first be deprotected, e.g. in the presence of a suitable acid, such as hydrochloric acid and the like, resulting in intermediates of formula (LXVI), followed by a reduction, resulting in a compound of formula (I-q-1) wherein Q represents H-Q₁, said compounds being represented by formula (I-q-1-1).

Intermediates of formula (IV), wherein G is ethyl substituted with hydroxy, said intermediates being represented by formula (IV-g-2) can also be prepared by reacting an intermediate of formula (XXXIII-a) with an intermediate of formula (LXVII) in the presence of a suitable base, such as sodium hydride, in a reaction-inert solvent, such as N,N-dimethylformamide.

A subgroup of intermediates of formula (IV-g-2), represented by formula (IV-g-2-1), can also be prepared by reacting an intermediate of formula (LXVIII) with an

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intermediate of formula (LXIX) in the presence of 1,3-dicyclohexylcarbodiimide, in a reaction-inert solvent, e.g. toluene.

Intermediates of formula (LXV) can be prepared by reacting an intermediate of formula (XXXIII-a) with an intermediate of formula (LXX), wherein W₈ is a suitable leaving group, such as a halo atom, e.g. bromo, in the presence of a suitable base, e.g. sodium hydride, in a reaction-inert solvent, e.g. N,N-dimethylformamide.

$$P = Q_{1} = \begin{bmatrix} R^{1} \\ Q_{2}(=0) \\ N \end{bmatrix} = \begin{bmatrix} R^{1} \\ Q_{2}($$

Intermediates of formula (V) can be prepared by reacting an intermediate of formula (LXXI) with 1*H*-isoindole-1,3 (2*H*)-dione in the presence of triphenylphosphine and diethyl azodicarboxylate.

Intermediates of formula (V) may also be prepared by reacting an intermediate of formula (LXXII) with 1*H*-isoindole-1,3 (2*H*)-dione in the presence of a suitable base, such as sodium hydride, and a suitable solvent, such as *N*, *N*-dimethylformamide.

$$Q_{2} = \begin{pmatrix} R^{1} & & & \\ Q_{2} & & & \\ Q_{3} & & & \\ Q_{4} & & & \\ Q_{3} & & & \\ Q_{4} & & & \\ Q_{5} & & & \\ Q_{5} & & & \\ Q_{6} & & & \\ Q_{7} & & & \\ Q_{8} & & \\ Q_{8} & & & \\ Q_{8} & & \\ Q_{8} & & \\ Q_{8} & & \\ Q_{8} & & \\$$

Intermediates of formula (LXXII) can be prepared by reacting an intermediate of formula (LXXII) with an intermediate of formula (LXXIII), wherein W₉ represents a suitable leaving group, such as a halo atom, e.g. chloro, in the presence of a suitable base, such as N, N -diethyl-ethanamine, and a suitable solvent, such as methylene chloride.

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Intermediates of formula (V), wherein in the definition of Q_2 , R^2 is C_{1-10} alkyl, said Q_2 being represented by C_{1-10} alkyl- Q_{1b} , and said intermediates by formula (V-a), can be prepared by reacting a compound of formula (I-a-3) with an intermediate of formula (LXXIV), wherein W_{10} is a suitable leaving group, such as a halo atom, e.g. chloro, in the presence of a suitable base, such as dipotassium carbonate, and a suitable solvent, such as acetonitrile.

$$H = Q_{1b} = \begin{pmatrix} R^1 \\ A^2 \\ A^3 \end{pmatrix} = \begin{pmatrix} R^1 \\ A^2 \\ A^3 \end{pmatrix}$$

$$(LXXIV) = \begin{pmatrix} R^1 \\ C_{1-10}alky \\ C_{1$$

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Intermediates of formula (LXXI) wherein, in the definition of Q₂, the carbon atom carrying the hydroxy, also carries two hydrogen atoms, said HO-Q₂ being represented by HO-CH₂-Q₂, and said intermediates being represented by formula (LXXI-a), can be prepared by reducing an intermediate of formula (LXXV) in the presence of a suitable reducing agent, such as lithium aluminium hydride, in a suitable reaction-inert solvent, e.g. tetrahydrofuran.

$$C_{1-4}$$
alkyl $-O-C(=O)-Q_2$
 N
 A_1
 A_2
 A_3
 A_4
 A_3
 A_4
 A_3
 A_4
 A_3
 A_4
 A_4
 A_3
 A_4
 A

Intermediates of formula (LXXI), wherein, in the definition of Q_2 , the carbon atom carrying the hydroxy, carries also at least one hydrogen, said HO- Q_2 being represented by HO- Q_3 H, and said intermediates being represented by formula (LXXI-b), can be prepared by reducing an intermediate of formula (IX) with a suitable reducing agent, e.g. sodium borohydride, in a reaction-inert solvent, e.g. an alcohol.

$$(O=)Q_3 \longrightarrow \begin{pmatrix} R^1 \\ N \\ A \end{pmatrix} = \begin{pmatrix} R^1 \\ A^2 \\ A \end{pmatrix} = \begin{pmatrix} R^1 \\ N \\ A \end{pmatrix} = \begin{pmatrix} R^1 \\ A \\ A \end{pmatrix}$$

Intermediates of formula (VI) wherein, in the definition of Q₂, R² is C₁₋₁₀alkyl substituted with N(P)₂ and the carbon atom adjacent to the nitrogen atom carrying the R² substituent carries also at least one hydrogen atom, said Q₂ being represented by (P)₂N-C₁₋₁₀alkyl-NH-Q_{2a}H, and said intermediates being represented by formula (VI-a), can be prepared by reductive amination of an intermediate of formula (LXXVI) with an intermediate of formula (LXXVII) in the presence of a suitable reductive agent, such as hydrogen, and a suitable catalyst, such as palladium-on-charcoal, platinum-on-charcoal, and the like, and optionally in the presence of a suitable catalyst poison, such as a thiophene solution. A suitable solvent in this reaction is a reaction-inert solvent, such as an alcohol.

$$(O=)Q_{2\overline{a}} = (D-1)Q_{2\overline{a}} = (D-1)Q_{\overline{a}} = (D-1)Q_{2\overline{a}} = (D-1)Q_{2\overline{a}} = (D-1)Q_{2\overline{a}} = (D-1)Q$$

Intermediates of formula (LXXVI) can be prepared by deprotecting an intermediate of formula (LXXVIII) in the presence of a suitable acid, such as hydrochloric acid and the like, in a suitable solvent, e.g. water.

$$(LXXVIII)$$

$$Q_{2a}$$

$$N$$

$$A^{1}$$

$$A^{2}$$

$$A^{2}$$

$$A^{3}$$

$$A^{2}$$

$$A^{3}$$

$$A^{$$

Intermediates of formula (IX) may be prepared by deprotecting an intermediate of formula (LXXIX) in the presence of a suitable acid, e.g. hydrochloric acid and the like.

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Intermediates of formula (LXXIX) can be prepared by reacting an intermediate of formula (LXXX) with an intermediate of formula (III) in the presence of a suitable base, e.g. dipotassium carbonate, in a suitable reaction-inert solvent, e.g. acetonitrile.

Intermediates of formula (LXXX) wherein, in the definition of Q_3 , the X^1 or X^2 moiety of the radicals of formula (b-1) to (b-8) represent NH, said Q_3 being represented by Q_3 -NH, and said intermediates being represented by formula (LXXX-a), may be prepared by cyclizing an intermediate of formula (LXXXI) in the presence of mercury oxide and sulphur, in a suitable reaction-inert solvent, e.g. an alcohol.

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Intermediates of formula (LXXXI) can be prepared by reducing an intermediate of formula (LXXXII) in the presence of a suitable reducing agent, such as hydrogen, in the presence of a suitable catalyst, such as palladium-on-charcoal, platinum-on-charcoal and the like, in a suitable solvent, e.g. a mixture of ammonia in alcohol. Suitable alcohols are methanol, ethanol, 2-propanol and the like.

Intermediates of formula (LXXXII) can be prepared by reacting an intermediate of formula (LXXXIII) with an intermediate of formula (LXXXIV) in a suitable reaction-inert solvent, e.g. ethanol.

Intermediates of formula (IX), wherein, in the definition of Q_3 , R^2 comprises C_{1-10} alkyl, said Q_3 being represented by C_{1-10} alkyl- Q_{1b} , and said intermediates being represented by formula (IX-a), can be prepared by reacting a compound of formula (I-a-3) with a reagent of formula (LXXXV), wherein (O=) C_{1-10} alkyl represents a carbonyl derivative of C_{1-10} alkyl and wherein W_{11} is a suitable leaving group, such as a halo atom, e.g. bromo, in a reaction-inert solvent, e.g. acetonitrile, in the presence of a suitable base, e.g. dipotassium carbonate.

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$$H-Q_{1b} \xrightarrow{N} \overset{a^1}{\underset{a^4=a^3}{\downarrow}} \overset{a^2}{\underset{a^4=a^3}{\downarrow}} + (O=)C_{1-10}alkyl-W_{11} \xrightarrow{\qquad} (O=)C_{1-10}alkyl-Q_{1b} \xrightarrow{N} \overset{a^1}{\underset{a^4=a^3}{\downarrow}} \overset{a^2}{\underset{a^4=a^3}{\downarrow}}$$

$$(I-a-3)$$

Intermediates of formula (X) wherein Q₄ comprises C₁₋₉alkyl, said Q₄ being represented by C₁₋₉alkyl-Q_{1b}, and said intermediates being represented by formula (X-a), can be prepared by reacting a compound of formula (I-a-3) with a reagent of formula (LXXXVI) wherein W₁₂ represents a suitable leaving group, such as a halo atom, e.g. chloro, in a reaction-inert solvent, e.g. 3-methyl-2-butanone, in the presence of a suitable base, e.g. dipotassium carbonate, sodium bicarbonate and the like.

$$H-Q_{1b} \xrightarrow{N} \stackrel{a^1}{\underset{a^4}{=}} \stackrel{a^2}{\underset{a^3}{=}} +W_{12}-C_{1-9alkyl}-CN \xrightarrow{NC-C_{1-9alkyl}} \stackrel{NC-C_{1-9alkyl}}{\underset{a^4}{=}} \stackrel{R^1}{\underset{a^3}{=}} \stackrel{A^2}{\underset{a^3}{=}} \stackrel{A^2}{\underset{a^3}{=}} \stackrel{A^2}{\underset{a^4}{=}} \stackrel{A^2}{\underset{a^3}{=}} \stackrel{A^2}{\underset{a^4}{=}} \stackrel{A^2}{\underset{a^3}{=}} \stackrel{A^2}{\underset{a^4}{=}} \stackrel{A^2}{\underset{a^3}{=}} \stackrel{A^2}{\underset{a^4}{=}} \stackrel{A^2}{\underset{a^3}{=}} \stackrel{A^2}{\underset{a^4}{=}} \stackrel{A^2}{\underset{a^4}$$

Intermediates of formula (X), wherein NC-Q₄ represents NC-(C_{1-9} alkyl)(R^4)N-C(=O)-Alk- X^1 , said intermediates being represented by formula (X-b), can be prepared by reacting an intermediate of formula (LXXXVII) with an intermediate of formula (LXXXVIII) in the presence of di-1*H*-imidazol-2-yl-methanone, a suitable base, such as N, N-diethyl-ethanamine, and a suitable solvent, such as methylene chloride.

HO-C-Alk-X¹
N
$$= \begin{bmatrix} R^1 \\ Alk-X^1 \end{bmatrix} + NC-C_1-9alkyl - NH$$

$$= \begin{bmatrix} R^4 \\ Alk-X^1 \end{bmatrix} + NC-C_1-9alkyl - NH$$

$$= \begin{bmatrix} R^4 \\ Alk-X^1 \end{bmatrix} + NC-C_1-9alkyl - N-C-Alk-X^1 + N-C-Alk-X^1 +$$

Intermediates of formula (XI), wherein Q₄ represents Q_{1b}, said intermediates being represented by formula (XI-a), can be prepared by reacting a compound of formula (I-a-3) with an intermediate of formula (LXXXIX), wherein W₁₃ represents a suitable leaving group, such as a halo atom, e.g. chloro, in the presence of a suitable base, such as disodium carbonate, and in the presence of a suitable solvent, such as 3-methyl-2-butanone.

Intermediates of formula (XIX) can be prepared by reacting an intermediate of formula (XC) with a suitable acid, such as hydrochloric acid.

- Pure stereochemically isomeric forms of the compounds of formula (I) may be obtained by the application of art-known procedures. Diastereomers may be separated by physical methods such as selective crystallization and chromatographic techniques, e.g., countercurrent distribution, liquid chromatography and the like.
- 10 The compounds of formula (I) as prepared in the hereinabove described processes are generally racemic mixtures of enantiomers which can be separated from one another following art-known resolution procedures. The racemic compounds of formula (I) which are sufficiently basic or acidic may be converted into the corresponding diastereomeric salt forms by reaction with a suitable chiral acid, respectively chiral base. Said diastereomeric 15 salt forms are subsequently separated, for example, by selective or fractional crystallization and the enantiomers are liberated therefrom by alkali or acid. An alternative manner of separating the enantiomeric forms of the compounds of formula (I) involves liquid chromatography, in particular liquid chromatography using a chiral stationary phase. Said pure stereochemically isomeric forms may also be derived from the corresponding pure 20 stereochemically isomeric forms of the appropriate starting materials, provided that the reaction occurs stereospecifically. Preferably if a specific stereoisomer is desired, said compound will be synthesized by stereospecific methods of preparation. These methods will advantageously employ enantiomerically pure starting materials.
- 25 The compounds of formula (I) show antiviral properties. Viral infections treatable using the compounds and methods of the present invention include those infections

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brought on by ortho- and paramyxoviruses and in particular by human and bovine respiratory syncytial virus (RSV).

The *in vitro* antiviral activity against RSV of the present compounds was tested in a test as described in the experimental part of the description, and may also be demonstrated in a virus yield reduction assay. The *in vivo* antiviral activity against RSV of the present compounds may be demonstrated in a test model using cotton rats as described in Wyde et al. (Antiviral Research (1998), 38, 31-42).

Due to their antiviral properties, particularly their anti-RSV properties, the compounds of formula (I) or any subgroup thereof, their prodrugs, N-oxides, addition salts, quaternary amines, metal complexes and stereochemically isomeric forms, are useful in the treatment of individuals experiencing a viral infection, particularly a RSV infection, and for the prophylaxis of these infections. In general, the compounds of the present invention may be useful in the treatment of warm-blooded animals infected with viruses, in particular the respiratory syncytial virus.

The compounds of the present invention or any subgroup thereof may therefore be used as medicines. Said use as a medicine or method of treatment comprises the systemic administration to viral infected subjects or to subjects susceptible to viral infections of an amount effective to combat the conditions associated with the viral infection, in particular the RSV infection.

The present invention also relates to the use of the present compounds or any subgroup thereof in the manufacture of a medicament for the treatment or the prevention of viral infections, particularly RSV infection.

The compounds of the present invention or any subgroup thereof may be formulated into various pharmaceutical forms for administration purposes. As appropriate compositions there may be cited all compositions usually employed for systemically administering drugs. To prepare the pharmaceutical compositions of this invention, an effective amount of the particular compound, optionally in addition salt form or metal complex, as the active ingredient is combined in intimate admixture with a pharmaceutically acceptable carrier, which carrier may take a wide variety of forms depending on the form of preparation desired for administration. These pharmaceutical compositions are desirable in unitary dosage form suitable, particularly, for administration orally, rectally, percutaneously, or by parenteral injection. For example, in preparing the compositions in oral dosage form, any of the usual pharmaceutical

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media may be employed such as, for example, water, glycols, oils, alcohols and the like in the case of oral liquid preparations such as suspensions, syrups, elixirs, emulsions and solutions; or solid carriers such as starches, sugars, kaolin, lubricants, binders, disintegrating agents and the like in the case of powders, pills, capsules, and tablets.

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- disintegrating agents and the like in the case of powders, pills, capsules, and tablets. Because of their ease in administration, tablets and capsules represent the most 5 advantageous oral dosage unit forms, in which case solid pharmaceutical carriers are obviously employed. For parenteral compositions, the carrier will usually comprise sterile water, at least in large part, though other ingredients, for example, to aid solubility, may be included. Injectable solutions, for example, may be prepared in which the carrier comprises saline solution, glucose solution or a mixture of saline and 10 glucose solution. Injectable suspensions may also be prepared in which case appropriate liquid carriers, suspending agents and the like may be employed. Also included are solid form preparations which are intended to be converted, shortly before use, to liquid form preparations. In the compositions suitable for percutaneous administration, the carrier optionally comprises a penetration enhancing agent and/or a 15 suitable wetting agent, optionally combined with suitable additives of any nature in minor proportions, which additives do not introduce a significant deleterious effect on
- The compounds of the present invention may also be administered via oral inhalation or insufflation by means of methods and formulations employed in the art for administration via this way. Thus, in general the compounds of the present invention may be administered to the lungs in the form of a solution, a suspension or a dry powder, a solution being preferred. Any system developed for the delivery of solutions, suspensions or dry powders via oral inhalation or insufflation are suitable for the administration of the present compounds.

the skin.

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Thus, the present invention also provides a pharmaceutical composition adapted for administration by inhalation or insufflation through the mouth comprising a compound of formula (I) and a pharmaceutically acceptable carrier. Preferably, the compounds of the present invention are administered via inhalation of a solution in nebulized or aerosolized doses.

It is especially advantageous to formulate the aforementioned pharmaceutical compositions in unit dosage form for ease of administration and uniformity of dosage. Unit dosage form as used herein refers to physically discrete units suitable as unitary dosages, each unit containing a predetermined quantity of active ingredient calculated to produce the desired therapeutic effect in association with the required pharmaceutical carrier. Examples of such unit dosage forms are tablets (including scored or coated

tablets), capsules, pills, suppositories, powder packets, wafers, injectable solutions or suspensions and the like, and segregated multiples thereof.

In general it is contemplated that an antivirally effective daily amount would be from 0.01 mg/kg to 500 mg/kg body weight, more preferably from 0.1 mg/kg to 50 mg/kg body weight. It may be appropriate to administer the required dose as two, three, four or more sub-doses at appropriate intervals throughout the day. Said sub-doses may be formulated as unit dosage forms, for example, containing 1 to 1000 mg, and in particular 5 to 200 mg of active ingredient per unit dosage form.

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The exact dosage and frequency of administration depends on the particular compound of formula (I) used, the particular condition being treated, the severity of the condition being treated, the age, weight, sex, extent of disorder and general physical condition of the particular patient as well as other medication the individual may be taking, as is well known to those skilled in the art. Furthermore, it is evident that said effective daily amount may be lowered or increased depending on the response of the treated subject and/or depending on the evaluation of the physician prescribing the compounds of the instant invention. The effective daily amount ranges mentioned hereinabove are therefore only guidelines.

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Also, the combination of another antiviral agent and a compound of formula (I) can be used as a medicine. Thus, the present invention also relates to a product containing (a) a compound of formula (I), and (b) another antiviral compound, as a combined preparation for simultaneous, separate or sequential use in antiviral treatment. The different drugs may be combined in a single preparation together with pharmaceutically acceptable carriers. For instance, the compounds of the present invention may be combined with interferon-beta or tumor necrosis factor-alpha in order to treat or prevent RSV infections.

The following examples are intended to illustrate the present invention.

Experimental part

Hereinafter, "DMF" is defined as N,N-dimethylformamide, "DIPE" is defined as disopropyl ether.

35 A. Preparation of the intermediate compounds

Example A1

a) Sodium methoxide (0.2 mol) was added to a mixture of N-(4-piperidinyl)-1H-benzimidazol-2-amine dihydrobromide (0.1 mol) in methanol (389ml), the mixture

was cooled on an ice bath and stirred for 2 hours.

Di-tert-butyldicarbonate (0.1mol) was added to a cooled mixture on an ice bath and then stirred for 18 hours at room temperature. The mixture was evaporated and suspended in water/DIPE. The residue was filtered off, washed with water/DIPE and dried. The residue was boiled up in CH₃OH, yielding 17.46g (55.2%) of 1,1-dimethylethyl 4-(1*H*-benzimidazol-2-ylamino)-1-piperidinecarboxylate; mp. 249.4°C (interm. 1).

b) A mixture of intermediate (1) (0.05 mol), 2-(chloromethyl)quinoline monohydrochloride (0.055 mol) and sodium carbonate (0.075 mol) in DMF (250ml)
10 was stirred at 55°C overnight. The solvent was evaporated. The residue was taken up in H₂O and CH₂Cl₂. The organic layer was separated, dried, filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH 97/3 and 95/5). The pure fractions were collected and the solvent was evaporated. The residue was suspended in DIPE, filtered off and dried, yielding 13.5g (59%) of 1,1-dimethylethyl 4-[[1-(quinolinylmethyl)-1*H*-benzimidazol-2-yl]amino]-1-piperidinecarboxylate (interm. 2).

Example A2

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- a) A mixture of 5,6,7,8-tetrahydro-2(1H)-quinoxalinone in phosphoryl chloride (200ml) was stirred and refluxed for 3 hours. The solvent was evaporated. The residue was taken up in ice and CH_2Cl_2 . The mixture was basified with NH_4OH . The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated, yielding 34g (86%) of 2-chloro-5,6,7,8-tetrahydroquinoxaline (interm. 3).
- b) A mixture of intermediate (3), 1-bromo-2,5-pyrolidinedione (0.116 mol) and dibenzoyl peroxide (1.3g) in tetrachloromethane (400ml) was stirred and refluxed for 35 minutes, brought to room temperature and then filtered. The reaction was carried out again using the same quantities. The residues were combined. The solvent was evaporated. The residue (60g) was purified by column chromatography over silica gel (eluent: cyclohexane/EtOAc 85/5; 15-35 μm). Two pure fractions were collected and their solvents were evaporated, yielding 25 g (43%) of (±)-5-bromo-2-chloro-5,6,7,8-tetrahydroquinoxaline (interm. 4) and 12 g (21%) of (±)-8-bromo-2-chloro-5,6,7,8-tetrahydroquinoxaline.
 - c) A dispersion of sodium hydride in mineral oil (60%) (0.0518 mol) was added portionwise at 5°C under N_2 flow to a mixture of intermediate (1) (0.0471 mol) in DMF (200ml). The mixture was stirred at 5°C/10°C for 1 hour. A solution of intermediate (4) (0.0565 mol) in DMF (50ml) was added dropwise. The mixture was stirred at room temperature for 3 hours and poured out into H_2O . The precipitate was

filtered off and taken up in CH2Cl2. The organic solution was dried (MgSO4), filtered and the solvent was evaporated. The residue (32g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/ NH₄OH 95/5/0.1; 20-45 μm). The pure fractions were collected and the solvent was evaporated, yielding 13.3g (58%) of (\pm) -1,1-dimethylethyl 4-[[1-(2-chloro-5,6,7,8-tetrahydro-5-quinoxalinyl)-1Hbenzimidazol-2-yl]amino]-1-piperidinecarboxylate (interm. 5).

Example A3

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- a) 2,3-Butanedione (0.0776 mol) was added at room temperature to a solution of sodium pyrosulfite (0.1 mol) in water (75ml). The mixture was heated to 70°C and then added to a solution of ethyl 2,3-diaminobenzoate (0.0776 mol) in water (75ml). 10 The mixture was stirred at 100°C for 12 hours, cooled, basified with K2CO3 10% and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (17.5g) was purified by column chromatography over silica gel (eluent: $CH_2Cl_2/EtOAc$ 93/7; 20-45 μm). The pure fractions
- 15 were collected and the solvent was evaporated, yielding 12g (67%) of ethyl 2,3-dimethyl-5-quinoxalinecarboxylate (interm. 6).
 - b) Lithium chloride (0.6 mol) was added portionwise at 80°C to a mixture of intermediate (6) (0.06 mol) and potassium tetrahydroborate (0.6 mol) in tetrahydrofuran (300ml). The mixture was stirred at 80°C for 5 hours, cooled, poured out into
- 20 H₂O and extracted with EtOAc. The organic layer was separated, washed with H₂O, dried (MgSO₄), filtered and the solvent was evaporated, yielding 10.5g (91%) of (±)-1,2,3,4-tetrahydro-2,3-dimethyl-5-quinoxaline-methanol (interm. 7).
 - c) MnO₂ (100g) was added portionwise at room temperature to a mixture of intermediate (7) (0.0546 mol) in dichloromethane (500ml). The mixture was stirred at room temperature overnight, filtered over celite, washed with CH2Cl2 and the filtrate was evaporated. The product was used without further purification, yielding 7.8g (77%) of 2,3-dimethyl-5-quinoxalinecarboxaldehyde (interm. 8).
 - d) Sodium tetrahydroborate (0.084 mol) was added portionwise at 5°C to a mixture of intermediate (8) (0.042 mol) in methanol (100ml). The mixture was stirred at 5°C for
- 30 30 minutes, hydrolized cold and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated, yielding 6.7g (85%) 2,3-dimethyl-5-quinoxalinemethanol (interm. 9).
- e) Thionyl chloride (0.045 mol) was added dropwise at 5°C to a mixture of intermediate (9) (0.03 mol) in dichloromethane (50ml). The mixture was stirred at 35 room temperature for 2 hours, poured out on ice and K2CO3 10%. The organic layer was separated, washed with K2CO3 10%, dried (MgSO4), filtered and the solvent was

evaporated. The product was used without further purification, yielding 6.2g (quant.) of 5-(chloromethyl)-2,3-dimethyl-quinoxaline (interm. 10).

f) A dispersion of sodium hydride in mineral oil (60%) (0.021 mol) was added portionwise at 5°C under N₂ flow to a mixture of intermediate (1) (0.02 mol) in DMF
5 (30ml). The mixture was stirred at 5°C under N₂ flow for 1 hour. A solution of intermediate (10) (0.03 mol) in a small amount of DMF was added dropwise at 5°C. The mixture was stirred at room temperature under N₂ flow for 2 hours, hydrolized and extracted with EtOAc. The organic layer was separated, washed several times with H₂O, dried (MgSO₄), filtered and the solvent was evaporated. The residue (12.5g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/ CH₃OH/NH₄OH 97.5/2.5/0.1; 20-45 μm). Two pure fractions were collected and their solvents were evaporated, yielding 7.8g (80%) of 1,1-dimethylethyl 4-[[1-[(2,3-dimethyl-5-quino-xalinyl)methyl]-1H-benzimidazol-2-yl]amino]-1-piperidinecarboxylate (interm. 11).

Example A4

8-Bromo-2-methylquinoline (0.0675 mol) was added portionwise at -70°C under N₂ flow to a mixture of a solution of butyllithium in hexane (1.6M) (0.135 mol) in tetrahydrofuran (300ml) and diethyl ether (300ml). The mixture was stirred for 30 minutes. A solution of DMF (0.405 mol) in tetrahydrofuran (100ml) was added quickly. The mixture was cooled to -70°C and stirred for 15 minutes. Ethanol (70ml) and a NH₄Cl solution 10% were added. The mixture was brought to room temperature and stirred for 15 minutes. NH₄Cl was added. The mixture was extracted with EtOAc. The organic layer was separated, washed with H₂O, dried (MgSO₄), filtered and the solvent was evaporated. The product was used without further purification, yielding 15g (>100%) of 2-methyl-8-quinolinecarboxaldehyde (interm. 12).

25 Example A5

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a) A mixture of 3-methoxy-2-methylquinoline (0.081 mol) in trifluoro-acetic acid (150ml) was hydrogenated at room temperature under a 3-4 bar pressure for 48 hours with palladium on activated carbon (2g) as a catalyst. After uptake of hydrogen (2 equiv.), the catalyst was filtered through celite and washed with H₂O. The filtrate was basified with a concentrated NH₄OH solution and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated, yielding 14.3g (quant.) of 5,6,7,8-tetrahydro-3-methoxy-2-methylquinoline (interm. 13). b) 3-Chlorobenzenecarboperoxoic acid (0.1 mol) was added portionwise at 5°C to a mixture of intermediate (13) (0.067 mol) in dichloromethane (300ml). The mixture was stirred at room temperature overnight, basified with K₂CO₃ 10% and separated into its layers. The aqueous layer was extracted with CH₂Cl₂. The combined organic

layer was dried (MgSO₄), filtered and the solvent was evaporated, yielding 13.7g (quant.) of 5,6,7,8-tetrahydro-3-methoxy-2-methylquinoline, 1-oxide (interm. 14).

- c) A mixture of intermediate (14) (0.067 mol) in acetic anhydride (100ml) was stirred at 90°C for 1 hour, poured out on ice and basified with NaOH 3N. CH₂Cl₂ was added.
- The organic layer was separated, washed with a diluted NaOH solution, dried (MgSO₄), filtered and the solvent was evaporated, yielding 16.8g (quant.) of 5,6,7,8-tetrahydro-3-methoxy-2-quinolinemethanol acetate (ester) (interm. 15).
 - d) A mixture of intermediate (15) (0.067 mol) and sodium hydroxide (13g) in methanol (60ml) was stirred and refluxed for 20 minutes, poured out on ice and
- extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated, yielding 12.3g (95%) of 5,6,7,8-tetrahydro-3-methoxy-2-quinolinemethanol (interm. 16).
 - In a similar way was also prepared (±)-5,6,7,8-tetrahydro-2-methyl-8-quinolinol (interm. 17).

15 Example A6

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Phosphorus tribromide (0.0105 mol) was added dropwise at $0^{\circ}\text{C}/5^{\circ}\text{C}$ under N_2 flow to a mixture of (±)-5,6,7,8-tetrahydro-2-methyl-8-quinolinol (intermediate 17) (0.03 mol) in toluene (20ml). The mixture was brought to room temperature and stirred at room temperature overnight. Ice water was added. The mixture was basified with a concentrated NaOH solution and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (6g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH 99/1;

20-45 μm). The pure fractions were collected and the solvent was evaporated, yielding 2g (29%) of (±)-8-bromo-5,6,7,8-tetrahydro-2-methylquinoline (interm. 18).

25 Example A7

- a) A mixture of N-2,6-dimetyl-2,3-pyridinediamine (0.122 mol) in trifluoro-acetic acid (250ml) was stirred and refluxed for 6 hours and brought to room temperature. The solvent was evaporated. The residue was taken up in CH₂Cl₂ and K₂CO₃ 10%. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated.
- The residue (32g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/EtOAc 97/3; 20-45 μm). The pure fractions were collected and the solvent was evaporated. The residue was taken up in petroleum ether. The precipitate was filtered off and dried, yielding 15g of residue (fraction 1). The mother layer was evaporated. The residue was combined with 14.1g of fraction 1, yielding 28.9 g of
- 35 1,6-dimethyl-2-(trifluoromethyl)-1*H*-imidazo[4,5-b]pyridine; mp. 100°C (interm. 19).

- b) 1-Bromo-2,5-pyrolidinedione (0.0735 mol) and dibenzoyl peroxide (1.5g) were added at room temperature to a solution of intermediate (19) (0.07 mol) in tetrachloromethane (450ml). The mixture was stirred and refluxed for 7 hours, then brought to room temperature and filtered. The reaction was carried out again using the same quantities. The mixtures were combined. The solvent was evaporated. The residue (50g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH 100/0 and 98/2; 20-45 μm). The pure fractions were collected and the solvent was evaporated, yielding 20.2g (49%) of 6-(bromomethyl)-1-methyl-2-(trifluoromethyl)-1-imidazo[4,5-b]pyridine (interm. 20).
- c) A mixture of ethyl 4-(1H-benzimidazol-2-ylamino)-1-piperidine-carboxylate (0.0464 mol), intermediate (20) (0.051 mol) and potassium carbonate (0.1392 mol) in acetonitrile (250ml) was stirred and refluxed for 90 minutes and then brought to room temperature. Water was added and the mixture was extracted twice with CH₂Cl₂. The combined organic layer was dried (MgSO₄), filtered and the solvent was evaporated.
- The product was used without further purification, yielding 23g (>100%) of ethyl 4[[1-[[1-methyl-2-(trifluoromethyl)-1*H*-imidazo[4,5-b]pyridin-6-yl]methyl]-1*H*benzimidazol-2-yl]amino]-1-piperidinecarboxylate (interm. 21).

Example A8

A mixture of ethyl 4-(1*H*-benzimidazol-2-ylamino)-1-piperidine-carboxylate (0.0289 mol), 7-chloro-6,7-dihydro-5*H*-cyclopenta[b]pyridine (0.0289 mol) and potassium carbonate (0.0867 mol) in acetonitrile (250ml) was stirred and refluxed for 48 hours and then brought to room temperature. The reaction was carried out again using the same quantities. The mixtures were combined, poured out into H₂O and extracted with EtOAc. The organic layer was separated, washed with H₂O, dried (MgSO₄), filtered and the solvent was evaporated. The residue (25g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/ CH₃OH/NH₄OH 97/3/0.5; 20-45 μm). Two fractions were collected and their solvents were evaporated, yielding 8g of ethyl 4-[[1-(6,7-dihydro-5*H*-1-pyrindin-7-yl)-1*H*-benzimidazol-2-yl]amino]-1-piperidinecarboxylate (interm. 22).

30 Example A9

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a) A dispersion of sodium hydride in mineral oil (0.261 mol) was added portionwise at room temperature under N_2 flow to a mixture of N-8-quinolinylformamide (0.174 mol) in DMF (500ml). The mixture was stirred at room temperature for 1 hour. A solution of 1-chloro-2-nitrobenzene (0.53 mol) in DMF (200ml) was added dropwise. The mixture was stirred at 140°C for 12 hours and then brought to room temperature. H_2O was added and the mixture was extracted with CH_2Cl_2 . The organic layer was

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separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (110g) was purified by column chromatography over silica gel (eluent: $CH_2Cl_2/cyclohexane$ 80/20; 20-45 μ m). The pure fractions were collected and the solvent was evaporated, yielding 9.8g (21%) of N-(2-nitrophenyl)-8-quinolinamine (interm. 23).

- b) A mixture of 6-quinolinemethanamine (0.074 mol), 2-chloro-3-nitropyridine (0.0888 mol) and potassium carbonate (0.185 mol) in acetronitrile (200ml) was stirred and refluxed for 5 hours and then cooled to room temperature. EtOAc and H₂O were added. The mixture was extracted with HCl 3N. The aqueous layer was basified with K₂CO₃ solid and extracted with CH₂Cl₂. The combined organic layer was dried
- 10 (MgSO₄), filtered and the solvent was evaporated, yielding 17.8g (84%) of N-(3-nitro-2-pyridinyl)-8-quinolinemethanamine (interm. 24).

Example A10

- a) A mixture of intermediate (24) (0.064 mol) in methanol (200ml) was hydrogenated under a 3 bar pressure for 2 hours with Raney nickel (10g) as a catalyst. After uptake
- of hydrogen (3 equiv), the catalyst was filtered through celite and the filtrate was evaporated, yielding 14.8g (93%) of N2-(8-quinolinylmethyl)-2,3-pyridinediamine (interm. 25).
 - b) A mixture of intermediate (25) (0.059 mol) and ethyl 4-isothiocyanato-1-piperidine-carboxylate (0.059 mol) in methanol (150ml) was stirred and refluxed for 4 hours and brought to room temperature. The solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: CH_2Cl_2/CH_3OH 97/3; 20-45 μ m). The desired fractions were collected and the solvent was evaporated, yielding 10.5g
 - (37%) of ethyl 4-[[[2-[(8-quinolinylmethyl)amino]-3-pyridinyl]amino]sulfinyl]-amino]-1-piperidine-carboxylate (interm. 26)
- c) A mixture of intermediate (26) (0.026 mol), mercury(II) oxide (0.052 mol) and sulfur (0.2g) in ethanol (120ml) was stirred and refluxed for 2 hours, brought to room temperature and filtered over celite. The filtrate was evaporated, yielding 8.7g (96%) of 4-[[1-(8-quinolinylmethyl)-1*H*-imidazo[4,5-b]pyridin-2-yl]amino]-1-piperidine-carboxylate (interm. 27).

30 Example A11

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- a) A mixture of 8-quinolinecarboxaldehyde (0.092 mol) and 4-methylbenzenesulfonic acid (0.3g) in 2-ethoxyethanol (110ml) was stirred and refluxed for 24 hours using a Dean Stark apparatus. The solvent was evaporated. The reaction was carried out again using the same quantities. The residues were combined and taken up in CH₂Cl₂. The
- organic solution was washed with K₂CO₃ 10% and decanted. The organic layer was dried (MgSO₄), filtered and the solvent was evaporated. The residue (41g) was

purified by column chromatography over silica gel (eluent: CH_2Cl_2/CH_3OH 98/2; 20-45 µm). Two pure fractions were collected and their solvents were evaporated, yielding 20g (34%) of 8-[bis(2-ethoxyethoxy)methyl]quinoline (interm. 28). b) A mixture of 8-quinolinecarboxaldehyde (0.248 mol), triethoxymethane (0.4464 mol) and 4-methylbenzenesulfonic acid (4g) in ethanol (250ml) was stirred and refluxed for 1 hour, brought to room temperature, poured out into K_2CO_3 10% and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The product was used without further purification, yielding 48.5g (80%) of 8-(diethoxymethyl)-quinoline (interm. 29).

c) A mixture of 2-quinolinecarboxaldehyde (0.08 mol) and 4-methylbenzenesulfonic acid (0.25g) in ethanol (100ml) was stirred and refluxed for 48 hours and brought to room temperature. The reaction was carried out again using the same quantities. The mixtures were combined. The solvent was evaporated. The residue was taken up in CH₂Cl₂. The organic solution was washed with K₂CO₃ 10% and with H₂O, then dried (MgSO₄), filtered and the solvent was evaporated. The product was used without further purification, yielding 32.5g of 2-(diethoxymethyl)quinoline (interm. 30).

Example A12

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Intermediate (1) (0.0377 mol) and intermediate (29) (0.0755 mol) were heated at 160°C for 1 hour and then purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH 98/2; 15-35 µm). The pure fractions were collected and the solvent was evaporated, yielding 15g (79%) of (±)-1,1-dimethylethyl 4-[[1-[ethoxy(8-quino-linyl)methyl]-1*H*-benzimidazol-2-yl]amino]-1-piperidinecarboxylate (interm. 31).

Example A13

4-Methylbenzenesulfonyl chloride (0.2222 mol) was added portionwise at 10°C to a
mixture of 1,1-dimethylethyl [1-(hydroxymethyl)-2-methylpropyl]carbamic acid (ester) (0.202 mol) in pyridine (65ml). The mixture was stirred at 10°C for 2 hours. H₂O (75ml) was added at 10°C. The precipitate was filtered off, washed with H₂O and taken up in CH₂Cl₂. The organic solution was washed with H₂O, dried, filtered and the solvent was evaporated, yielding 49g (68%) of (±)-1,1-dimethylethyl [1-[[[(4-30 methylphenyl)sulfonyl]oxy]methyl]-2-methylpropyl]carbamate; mp. 85°C(interm. 32).

Example A14

a) A mixture of compound (33) (0.0347 mol), 1-bromo-3-methyl-2-butanone (0.052 mol) and potassium carbonate (0.104 mol) in acetonitrile (255ml) was stirred and refluxed for 2 hours and filtered. The filtrate was evaporated. The residue was taken up in H_2O and the mixture was extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The product was

used without further purification, yielding 16.84g of (±)-1-[4-[[1-[ethoxy(8-quinolinyl)methyl]-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]-3-methyl-2-butanone (interm. 34) (quant.).

In a similar way were also prepared:

- 5 1-[4-(1*H*-benzimidazol-2-ylamino)-1-piperidinyl]-3-methyl-2-butanone; 1-[4-[[1-(8-quinolinyl)-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]-3-methyl-2-butanone; and 1-[4-[[1-(2-quinolinylmethyl)-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]-3-methyl-2-butanone.
- b) A mixture of intermediate (34) (0.036 mol) in methanol (200ml) was stirred at 10°C. Sodium tetrahydroborate (0.04 mol) was added portionwise. The mixture was stirred for 90 minutes. H₂O was added. The solvent was evaporated. The residue was extracted with CH₂Cl₂. The organic layer was separated, washed with H₂O, dried (MgSO₄), filtered and the solvent was evaporated, yielding 17g (96%) of (±)-4-[[1-
- 15 [ethoxy(8-quinolinyl)methyl]-1*H*-benzimidazol-2-yl]amino]-alpha-(1-methylethyl)-1-piperidineethanol (interm. 35).
 - c) Diethyl azodicarboxylate (0.015 mol) was added dropwise at 0°C under N₂ flow to a solution of intermediate (35) (0.01 mol), phthalimide (0.015 mol) and triphenyl-phosphine (0.015 mol) in tetrahydrofuran (100ml). The mixture was stirred at room
- temperature for 2 hours. EtOAc was added. The mixture was extracted with HCl 3N and separated into its layers. The aqueous layer was washed twice with EtOAc, basified with K₂CO₃ solid and extracted with CH₂Cl₂. The combined organic layer was dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH 97/3/0.2;
- 20-45 μm). Two pure fractions were collected and their solvents were evaporated, yielding 2.3g (30%) of (±)-2-[2-[4-[[1-[ethoxy(8-quinolinyl)methyl]-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]-3-methylbutyl]-1*H*-isoindole-1,3(2*H*)dione (interm.

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and Et₃N (0.072 mol) in CH₂Cl₂ (100ml) was cooled to 0°C under N₂ flow. A mixture of methanesulfonyl chloride (0.036 mol) in CH₂Cl₂ (a small amount) was added dropwise. The mixture was allowed to cool to room temperature while stirring for 3 hours. Water was added. The mixture was decanted. The organic layer was dried (MgSO₄), filtered and the solvent was evaporated, yielding 8.5g of intermediate (80) (86%).

e) Preparation of intermediate

A solution of 1*H*-isoindole-1,3(2*H*)-dione (0.0828 mol) in DMF (80ml) was cooled to 10°C. NaH 60% in oil (0.0828 mol) was added portionwise. The mixture was allowed to cool to room temperature while stirring for 1 hour. A mixture of intermediate (80) (0.0207 mol) (prepared according to A14d) in DMF (a small amount) was added dropwise. The mixture was stirred at room temperature for 1.5 hours, at 60°C for 5 hours and at room temperature for the weekend. The residue (9.6g) was crystallized from diethyl ether and CH₃CN. The precipitate was filtered off and dried, yielding 4g of intermediate (81) (42%).

Example A15

a) A mixture of 1-[4-(1*H*-benzimidazol-2-ylamino)-1-piperidinyl]-3-methyl-2-butanone (0.03 mol) and benzenemethanamine (0.09 mol) in methanol (200ml) was hydrogenated at 40°C under a 3 bar pressure for 48 hours with palladium on activated carbon (1.3g) as a catalyst. After uptake of hydrogen, the catalyst was filtered through celite, washed with CH₃OH and the filtrate was evaporated. Hydrogenation was continued for 24 hours. After uptake of hydrogen, the catalyst was filtered through celite, washed with CH₃OH and the filtrate was evaporated. The residue was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH 85/14/1; 20-45 um). The desired fractions were cellected and the column through the catalyst.

- 25 20-45 μm). The desired fractions were collected and the solvent was evaporated. The residue was crystallized from diethyl ether. The precipitate was filtered off and dried, yielding 0.4g of (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1H-benzimidazol-2-amine; mp. 138°C (interm. 37).
- b) Di-tert-butyl dicarbonate (0.02 mol) was added at 5°C to a mixture of intermediate
 30 (37) (0.0186 mol) in dichloromethane (60ml). The mixture was stirred at room temperature for 3 hours and poured out into H₂O. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The product was used without

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further purification, yielding 5.9g of (±)-1,1-dimethylethyl [1-[[4-[[1-[(1,1-dimethylethoxy)carbonyl]-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]methyl]-2-methyl-propyl]carbamate (interm. 38).

Example A16

- A mixture of 1-[4-[[1-(8-quinolinyl)-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]3-methyl-2-butanone (0.0222 mol) and benzenemethanamine (0.0666 mol) in methanol (250ml) was hydrogenated at 40°C under a 3 bar pressure for 24 hours with palladium on activated carbon (1.5g) as a catalyst. After uptake of hydrogen, the catalyst was filtered through celite, washed with CH₂Cl₂ and CH₃OH and the filtrate was
- evaporated. Palladium on activated carbon (1.5g) and methanol (250ml) were added again. Hydrogenation was continued at 40°C under a 3 bar pressure for 24 hours. After uptake of hydrogen, the catalyst was filtered through celite, washed with CH₂Cl₂ and the filtrate was evaporated. The residue (22g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/ NH₄OH 95/5/0.1 and
- 85/15/1; 20-45 μm). Three pure fractions were collected and their solvents were evaporated, yielding 2.6g 1-[4-[[1-(1,2,3,4-tetrahydro-8-quinolinyl)-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]-3-methyl-2-butanone (interm. 40) (fraction 1), 2.9g of fraction 2 and 0.7g of fraction 3. Fraction 2 and 3 were crystallized from CH₃CN. The precipitate was filtered off and dried, yielding 0.82g (±)-N-[1-[3-methyl-2-
- [(phenylmethyl)amino]butyl]-4-piperidinyl]-1-(1,2,3,4-tetrahydro-8-quinolinyl)-1*H*-benzimidazol-2-amine; mp. 126°C and 0.55g of *N*-(4-piperidinyl)-1-(1,2,3,4-tetrahydro-8-quinolinyl)-1*H*-benzimidazol-2-amine; mp. 205°C (comp. 48).

Example A17

- a) A mixture of N-(4-piperidinyl)-1-(4-quinolinylmethyl)-1H-benzimidazol-2-amine (comp. 23) (0.0129 mol), chloroacetonitrile (0.0155 mol), potassium iodide (0.00129 mol) and potassium carbonate (0.0258 mol) in 4-methyl-2-pentanone (80ml) was stirred and refluxed for 5 hours. H₂O was added. The solvent was evaporated. H₂O and CH₂Cl₂ were added. The precipitate was filtered off. The filtrate was separated into its layers. The organic layer was dried (MgSO₄), filtered and the solvent was
- evaporated. The residue (3.5g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/ CH₃OH/NH₄OH 95/5/0.3; 15-40 μm). The pure fractions were collected and the solvent was evaporated. The residue was crystallized from CH₃CN. The precipitate was filtered off and dried, yielding 0.94g 4-[[1-(4-quinolinylmethyl)-1*H*-benzimidazol-2-yl]amino]-1-piperidineacetonitrile; mp. 190°C (interm. 41).
- b) A mixture of N-(4-piperidinyl)-[1,2'-bi-1H-benzimidazol]-2-amine (comp. 71) (0.01 mol), chloroacetonitrile (0.01 mol) and sodium hydrogen carbonate (0.02 mol) in

DMF (50ml) was stirred at 50°C overnight. The solvent was evaporated. The residue was taken up in H₂O and the mixture was extracted with CH₂Cl₂. The organic layer was separated, dried, filtered and the solvent was evaporated. The residue was suspended in DIPE, filtered off and dried, yielding 2.3g (63%) of product. This fraction was purified over silica gel on a glass filter (eluent: CH₂Cl₂/(CH₃OH/NH₃) 97/3). The pure fractions were collected and the solvent was evaporated, yielding 1.36g (37%) of 4-[(1,2'-bi-1H-benzimidazol-2-yl)amino]-1-piperidine-acetonitrile (interm. 42).

Example A18

Preparation of intermediate

- A mixture of 2-chloro-1*H*-benzimidazole (0.0189 mol) and 1,1-dimethylethyl 2-aminocyclohexanecarbamoate (0.04725 mol) (prepared according to A1a))was stirred at 140°C for 3 hours, then brought to room temperature and taken up in CH₂Cl₂/CH₃OH. The same procedure was repeated 3 times on the same quantities of 2-chloro-1*H*-benzimidazole and 1,1-dimethylethyl 2-aminocyclohexanecarbamoate.
- The mother layers were brought together, dried (MgSO₄), filtered and the solvent was evaporated. The residue (28g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/ NH₄OH 96/4/0.1; 15-35μm). Two fractions were collected and the solvent was evaporated, yielding 4.5g of intermediate (84) (24%).

Example A19

Preparation of intermediate

A mixture of quantities of ethyl 4-(1*H*-benzimidazol-2-ylamino)-1-piperidine-

according to A14d) and K₂CO₃ (0.0463 mol) in CH₃CN (50ml) and DMF (5ml) was stirred and refluxed for 6 hours, poured out into H₂O and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent:

25 CH₂Cl₂/CH₃OH 97/3; 35-70μm). The pure fractions were collected and the solvent was evaporated, yielding: 0.87g of intermediate (76) (13%).

Example A20

a) Preparation of intermediate

A solution of

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A1b) in HCl 6N (60ml) was stirred and refluxed for 12 hours and then brought to room temperature. The solvent was evaporated. The residue was taken up in 2-propanol. The precipitate was filtered off, washed with CH₃CN, washed with diethyl ether and dried, yielding: 4g of intermediate (82) (94%).

b) Preparation of intermediate

Intermediate (82 (0.0094 mol) was added at room temperature to CH₂Cl₂ (70ml). Et₃N (0.0188 mol) was added. 1,1'-carbonylbis-1*H*-imidazole (0.0188 mol) was added. The mixture was stirred at room temperature for 4.5 hours. (Methylamino)acetonitrile .HCl (0.0188 mol) was added. The mixture was stirred at room temperature for 12 hours. The organic layer was separated, washed twice with water, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH 98.5/1.5; 35-70 µm). The pure fractions were collected and the solvent was evaporated. The residue (2.2g) was crystallized from CH₃CN. The precipitate was filtered off and dried, yielding: 1.5g of intermediate (83) (41%).

Example A21

A mixture of intermediate

(prepared according to A1b) in HCl 3 N (200ml) was stirred and refluxed for 1 hour. The solvent was evaporated. The residue was taken up in EtOAc and NH4OH. The mixture was stirred for 30 minutes and filtered. The solvent was evaporated. The product was used

without further purification, yielding 14g of

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Tables 1, 2 and 3 list intermediates which were prepared analogous to one of the above examples.

Table 1

Int. No.	Ex. No.	Rª	R ^b	R°	n	a	*	b	R ^d	Re	R ^f	R ^g
43	A10c	Н	Н	Н	1	N	2	С	-	Н	Н	Н
44	A12	CH ₃	H	O(CH ₂) ₂ OC ₂ H ₅	1	СН	8	С	н	Н	Н	_
45	A12	CH₃	H	O(CH ₂) ₂ OC ₂ H ₅	1	СН	2	С	-	Н	н	Н
46	A7c	CH₃	H	Н	1	CH	2	N	-	OCH ₃	-	н
47	A7c	H	H	н -	1	СН	2	С	-	Н	н	CI
48	A7c	Н	H	Н	1	СН	2	С	-	Н	Cl	н
49	A7c	H	Н	н	1	CH	2	С	-	Н	Н	Н
2	A1b	CH ₃	H	Н	1	СН	2	С	-	Н	Н	н
50	A12	CH₃	CH ₃	OC₂H₅	1	СН	8	С	Н	Н	H	-
51	A12	CH ₃	Н	OC₂H₅	1	CH	2	C	-	Н	Н	н
52	A12	CH ₃	Н	OC ₂ H ₅	1	СН	2	C	-	OCH ₃	Н	Н
31	A12	CH ₃	Н	OC ₂ H ₅	1	СН	8	C	Н	Н	Н	_
53	A3f	н	Н	Н	1	СН	8	C	Н	Н	Н	-
54	A3f	CH ₃	Н	Н	1	СН	8	N	H	Н	-	

Int. No.	Ex. No.	Rª	R ^b	R ^c	n	а	*	b	R ^d	Re	R ^f	R ^g
55	A7c	CH ₃	Н	Н	1	CH	8	С	CH ₃	Н	Н	-
11	A3f	CH ₃	Н	Н	1	СН	8	N	CH ₃	CH₃	-	-
56	A7c	H	Н	Н	1	СН	4	С	H	Н	_	Н
57	A7c	Н	CH ₃	H	1	СН	8	С	Н	Н	H	-
27	A10c	Н	H	H	1	N	8	c	Н	н	Н	-
58	A10c	H	H	-	0	СН	8	С	Н	H	Н	-
66	A12	CH₃	CH ₃	O(C ₂ H ₅)OC ₂ H ₅	1	CH	8	С	Н	н	Н	-
67	A12	CH ₃	H	O(C ₂ H ₅)OC ₂ H ₅	1	CH	8	С	Н	H	н	-
68	A1b	CH ₃	CH ₃	CH ₃	1	CH	8	С	н	Н	Н	-
69	A1b	CH ₃	Н	H	1	СН	2	С	-	OCH₃	H	н
70	A1b	CH₃	H	H	1	СН	2	N	-	Н	-	н
71	A1b	CH ₃	Н	H	1	СН	8	С	OCH ₃	Н	H	-

^{* =} position bicyclic heterocycle

Table 2

Int. No.	Ex. No.	Rª	R ^b	n	L
59	A2c	СН3	H	0	N Ca
60	A8	Н	H	0	
61	A2c	н	Н	0	
5	A2c	СН3	Н	0	N CI
21	A7c	Н	Н	1	N CF_3 CH_3

			_ h	Γ-	I .
Int.	Ex. No.	Rª	R ^b	n	L
110.	1.10.	_		_	N.
62	A3f	CH ₃	Н	1	ОСН3
63	A7c	CH ₃	н	1	N
64	A7c	Н	Н	1	
04	1,0	**	**	1	S
65	A2c	CH ₃	Н	0	
		,		ľ	0
22	A8	н	Н	0	
					N
					N CI
72	A2c	CH ₃	CH ₃	0	
73	A2c	СН₃	CH₂	0	
					N CI
					N
74	A2c	CH ₃	CH ₃	0	
					, N. Ci
75	A2c	CH ₃	CH-	0	N CH ₃
		U113	C113		N CH ₃
					N N
76	A19	H	Н	1	

Table 3

1	Ex. No.	L	Physical data
77	Alb	NH O C(CH ₃) ₃	
78	А1ь	, о с(сн ₃) ₃	
79	Alb	H C(CH ₃) ₃	trans
80	A14d	H ₃ C CH ₃	
81	A14e	СН,	·
82	A20	но	
83	A20	CH ₃	

B. Preparation of the final compounds

Example B1

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- a) A mixture of 2-propanol and hydrochloric acid (15ml) was added to a mixture of intermediate (2) (0.0284 mol) in 2-propanol (150ml). The mixture was stirred and refluxed for 90 minutes and cooled. The precipitate was filtered off, washed with 2-propanol and DIPE and dried, yielding 10.36g of N-(4-piperidinyl)-1-(2-quinolinyl-methyl)-1H-benzimidazol-2-amine dihydrochloride (comp.1).
- b) A mixture of compound (1) (0.01 mol) and sodium carbonate (0.03 mol) in 4-methyl-2-pentanone (250ml) was stirred and refluxed for a few hours using a water separator (until gas development stops). 2-Bromoethyl carbamic acid 1,1-dimethylethyl ester (0.015 mol) was added. The mixture was stirred and refluxed for 18 hours using a water separator, then cooled, washed with H₂O, dried, filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: CH₂Cl₂/C₂H₅OH 95/5 and 90/10). The pure fractions were collected and the solvent was evaporated, yielding 3.8g of 1,1-dimethylethyl [2-[4-[[1-(2-quino-linylmethyl)-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]ethyl]carbamate (comp. 2). c) A mixture of compound (2) (0.0076 mol) in a mixture of 2-propanol and
 - hydrochloric acid (10ml) and 2-propanol (100ml) was stirred and refluxed for 1 hour

and then cooled. The precipitate was filtered off, washed with 2-propanol and DIPE and dried, yielding 3.08g of N-[1-(2-aminoethyl)-4-piperidinyl]-1-(2-quinolinyl-methyl)-1H-benzimidazol-2-amine tetrahydrochloride monohydrate (comp. 3).

- d) A mixture of compound (115) (0.00305 mol) in HBr/HOAc 33% (34ml) was stirred at room temperature for 2 hours, poured out on ice, basified with a concentrated NH₄OH solution and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH 96/4/0.2; 15-40 μm). Two fractions (F1 and F2) were collected and their solvents were evaporated, yielding
- 10 0.56g F1 (46%) and 0.69g F2 (50%). F1 was crystallized from diethyl ether. The precipitate was filtered off and dried, yielding 0.27g of (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-4-methyl-1-(8-quinolinylmethyl)-1H-benzimidazol-2-amine (comp. 116).
- e) A mixture of compound (155) (0.0024 mol) in CH₃OH (3ml) and 2-propanol (15ml)
 was stirred and refluxed for 2 hours, filtered, washed with 2-propanol and dried. The residue (1.05g) was taken up in CH₂Cl₂ and basified with NH₄OH. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (0.42g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH 85/15/2; 15-40 μm). The pure fractions were collected and the solvent was evaporated. The residue (0.35g) was dissolved in CH₃OH and converted into the
 - was evaporated. The residue (0.35g) was dissolved in CH₃OH and converted into the ethanedioic acid salt. The precipitate was filtered off and dried. This fraction was taken up in water and CH₂Cl₂ and alkalized with K₂CO₃ 10%. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (0.21g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/
- NH₄OH 75/28/1; 15-40 μ m). The pure fractions were collected and the solvent was evaporated, yielding 0.13g of compound (156).

Example B2

A mixture of intermediate (27) (0.02 mol) in hydrochloric acid (6N) (85ml) was stirred and refluxed at 50°C overnight and then brought to room temperature. The solvent was evaporated. The residue was taken up in K₂CO₃ 10% and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated, yielding 5g (69%) of N-(4-piperidinyl)-3-(8-quinolinylmethyl)-3H-imidazo[4,5-b]pyridin-2-amine (comp. 41).

Example B3

A mixture of intermediate (41) (0.00668 mol) in a solution of ammonia in methanol (7N) (70ml) was hydrogenated at room temperature under a 3 bar pressure for 5 hours

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with Raney nickel (2.7g) as a catalyst. After uptake of hydrogen (2 equiv.), the catalyst was filtered through celite, washed with CH₂Cl₂ and CH₃OH and the filtrate was evaporated. The residue was taken up in CH₂Cl₂ and a small amount of CH₃OH. The organic solution was washed with H₂O, dried (MgSO₄), filtered and the solvent was evaporated. The residue was crystallized from EtOAc. The precipitate was filtered off and dried, yielding 1.6g (60%) of N-[1-(2-aminoethyl)-4-piperidinyl]-1-(4-quinolinyl-methyl)-1H-benzimidazol-2-amine; mp. 196°C (comp. 24).

Example B4

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A mixture of intermediate (36) (0.00351 mol) in hydrazine (2.5ml) and ethanol (30ml) was stirred and refluxed for 20 minutes and brought to room temperature. Ice water was added. The mixture was extracted with CH₂Cl₂ and separated into its layers. The aqueous layer was washed twice with CH₂Cl₂. The combined organic layer was dried (MgSO₄), filtered and the solvent was evaporated. The residue was taken up in diethyl ether. The precipitate was filtered off and dried, yielding 1g of (±)-N-[1-[1-(aminomethyl)-2-methylpropyl]-4-piperidinyl]-1-[ethoxy(8-quinolinyl)methyl]-1H-benzimidazol-2-amine; mp. 202°C (comp. 100).

Example B5

Intermediate (32) (0.1382 mol) was added at 55°C to a mixture of (±)-1-[ethoxy(3-methoxy-2-quinolinyl)methyl]-N-(4-piperidinyl)-1H-benzimidazol-2-amine (0.0346 mol) and potassium carbonate (0.242 mol) in acetonitrile (108ml) and DMF (20ml) (1 equiv of intermediate (32) was added every hour). The mixture was stirred at 55°C for 1 hour and filtered. The filtrate was poured out into H₂O and the mixture was extracted with EtOAc. The organic layer was separated, washed with a saturated NaCl solution, dried (MgSO₄), filtered and the solvent was evaporated. The residue was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/ NH₄OH 98/2/0.4 and 96/4/0.5; 20-45 μm). Two fractions were collected and their solvents were evaporated, yielding 2.5g (23%) of (±)-1,1-dimethylethyl [1-[[4-[[1-[ethoxy(3-methoxy-2-quinolinyl)methyl]-1H-benzimidazol-2-yl]amino]-1-piperidinyl]methyl]-2-methylpropyl]carbamate (comp. 38).

30 Example B6

A mixture of 1-[4-[[1-(2-quinolinylmethyl)-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]-3-methyl-2-butanone (0.0158 mol) and benzenemethanamine (0.0474 mol) in methanol (150ml) was hydrogenated at 40°C under a 3 bar pressure for 48 hours with palladium on activated carbon (0.7g) as a catalyst. After uptake of hydrogen (1 equiv), the catalyst was filtered through celite, washed with CH₂Cl₂/ CH₃OH and the filtrate was evaporated. The residue (11.5g) was purified by column chromatography

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over silica gel (eluent: CH2Cl2/CH3OH/NH4OH 94/6/0.5; 20-45 µm). The pure fractions were collected and the solvent was evaporated, yielding 4g of residue. This fraction was converted into the hydrochloric acid salt with 2-propanol/ HCl. The precipitate was filtered off and dried, yielding 5.1g of product. This fraction was 5 converted into the free base and then purified by column chromatography over C₁₈ (eluent: CH₃OH/NH₄OAc 60/40 and 80/20; column: KROMASIL C18). Two pure fractions were collected and their solvents were evaporated, yielding 0.8g of fraction 1 and 2g of fraction 2. Fraction 1 was crystallized from diethyl ether. The precipitate was filtered off and dried, yielding 0.5g of (±)-N-[1-(2-amino-3-methylbutyl)-4-10 piperidinyl]-1-(2-quinolinylmethyl)-1H-benzimidazol-2-amine; mp. 135°C (comp. 6). Fraction 2 was dissolved in 2-propanol and converted into the hydrochloric acid salt (1:4). The precipitate was filtered off and dried, yielding 2.2g of $(\pm)-N-[1-(2-amino-3-1)]$ methylbutyl)-4-piperidinyl]-1-[(1,2,3,4-tetrahydro-2-quinolinyl)methyl]-1Hbenzimidazol-2-amine tetrahydrochloride monohydrate; mp. 230°C (comp. 46).

15 Example B7

- a) A dispersion of sodium hydride in a mineral oil (60%) (0.01 mol) was added portionwise at 0°C under N₂ flow to a mixture of intermediate (38) (0.005 mol) in DMF (25ml). The mixture was stirred at room temperature for 1 hour. A solution of 2-(bromomethyl)-3-methoxyquinoline (0.0055 mol) in DMF (10ml) was added dropwise. The mixture was stirred at room temperature for 2 hours had been declared at the contraction of the point of the contraction of the point of the contraction of th
- dropwise. The mixture was stirred at room temperature for 2 hours, hydrolized with K₂CO₃ 10% and extracted with EtOAc. The organic layer was separated, washed with NaCl, dried (MgSO₄), filtered and the solvent was evaporated, yielding 4.5g (>100%) of (±)-1,1-dimethylethyl [1-[[4-[[1-[(3-methoxy-2-quinolinyl)methyl]-1*H*-benzimi-dazol-2-yl]-amino]-1-piperidinyl]methyl]-2-methylpropyl]carbamate (comp. 14).
- b) A dispersion of sodium hydride in a mineral oil (60%) (0.014 mol) was added portionwise at 0°C under N₂ flow to a mixture of intermediate (38) (0.007 mol) in DMF (30ml). The mixture was stirred at 5°C for 1 hour. A solution of (±)-2,8-di-bromo-5,6,7,8-tetrahydroquinoline (0.0084 mol) in DMF (10ml) was added dropwise. The mixture was stirred at room temperature for 2 hours. H₂O and EtOAc were added.
- The organic layer was separated, washed with a saturated NaCl solution, dried (MgSO₄), filtered and the solvent was evaporated. The residue (5.6g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH 97/3/0.5; 20-45 μm). The pure fractions were collected and the solvent was evaporated, yielding 1.1g (25%) of (±)-1,1-dimethylethyl [1-[[4-[[1-(2-bromo-5,6,7,8-tetrahydro-8-35]]]]. The pure fractional 2 collected and the solvent was evaporated.
- quinolinyl)-1*H*-benzimidazol-2-yl]amino]-1-piperidinyl]methyl]-2-methylpropyl]-carbamate (comp. 55).

- c) A mixture of intermediate 84 (0.0145 mol), 8-bromomethylquinoline (0.0174 mol) and K₂CO₃ (0.029 mol) in CH₃N (70ml) was stirred and refluxed for 4 hours, then brought to room temperature. The solvent was evaporated. The residue was taken up in H₂O and extracted twice with CH₂Cl₂. The organic layer was separated, dried (MgSO₄),
- filtered and the solvent was evaporated. The residue was crystallized from diethyl ether/CH₃CN. The precipitate was filtered off and dried, yielding 5.07g of compound 79 (74%).

Example B8

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- c) (\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-[(5,6,7,8-tetrahydro-3-methoxy-2-quinolinyl)methyl]-1H-benzimidazol-2-amine tetrahydrochloride monohydrate (0.00218 mol) was basified with K₂CO₃ 10%. The mixture was extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was
- evaporated, to give A'. A mixture of A' in dichloromethane (50ml) was cooled to 0°C. A solution of tribromoborane in dichloromethane (0.01526 mol) was added dropwise.
- The mixture was stirred at room temperature overnight, poured out on ice, basified with a concentrated NH₄OH solution, decanted and extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (1.1g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH 90/10/0.5; 20-45 μm). The desired fractions were collected
- and the solvent was evaporated. The residue was converted into the hydrochloric acid salt (1:4) with HCl/2-propanol. The precipitate was filtered off and dried, yielding 0.5g (37%) of (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-[(5,6,7,8-tetrahydro-3-hydroxy-2-quinolinyl)methyl]-1H-benzimidazol-2-amine tetrahydro-chloride monohydrate; mp. 240°C (comp. 63).

25 Example B9

- a) A mixture of compound 158 (0.0089 mol) in HCl 3N (40ml) was stirred at 100°C for 12 hours, then brought to room temperature and poured out on ice and NH₄OH. EtOAc was added. The precipitate was filtered off, washed with EtOAc and dried, yielding 2g of compound 159.
- b) A mixture of compound 168 (0.00895 mol) in HCl 3N (35ml) was stirred at 100°C for 24 hours. The solvent was evaporated. The residue was taken up in EtOAc. The mixture was basified with NH₄OH. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. Part of this fraction (0.7g) was crystallized from CH₃CN. The precipitate was filtered off and dried, yielding 0.3g of compound
- 35 167.

c) A mixture of compound 176 (0.00373 mol) in HCl 3N (20ml) was stirred at 100°C for 12 hours, brought to room temperature, poured out on ice, basified with NH₄OH and extracted with EtOAc. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. This fraction was dissolved in 2-propanol and converted into the hydrochloric acid salt (1:3). The precipitate was filtered off and dried, yielding 1.5g of compound 173 (77%).

Example B10

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A mixture of intermediate

(prepared according to A1b)), 1,2-ethanediamine (0.02 mol) and NaCN (0.0002 mol) in CH₃OH (7ml) was heated at 45°C for 4 hours and then brought to room temperature. Water was added. The mixture was extracted with CH₂Cl₂. The organic layer was separated, dried (MgSO₄), filtered and the solvent was evaporated. The residue (0.65g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH 90/10/1; 35-70 μ m). The pure fractions were collected and the solvent was evaporated, yielding 0.42g of compound 170 (56%)

15 Example B11

A mixture of intermediate

(prepared according to A14a)) and formic acid/NH₃ (0.0462 mol) in formamide (35ml) was stirred at 140°C for 30 min and then brought to room temperature. CH_2Cl_2 was added. The organic layer was separated, washed with K_2CO_3 10%, dried (MgSO₄), filtered and the solvent was evaporated. The residue (4g) was purified by column chromatography over silica gel (eluent: $CH_2Cl_2/CH_3OH/NH_4OH$ 97/3/0.1; 15-40 µm). Two pure fractions were collected and their solvents were evaporated. The second fraction was crystallized from CH_3CN and diethyl ether. The precipitate was filtered off and dried, yielding: 1.37g of compound 137 (46%).

Example B12

Isopropyl titanate (IV) (0.0294 mol) was added at room temperature to a mixture of intermediate 85 (0.0245 mol) and 1-acetylpiperazine (0.027 mol) in CH₂Cl₂ (50ml) and ethanol (50ml). The mixture was stirred at room temperature for 7 hours. NaBH₃CN (0.0245 mol) was added portionwise. The mixture was stirred at room temperature for 12 hours. H₂O was added. The mixture was filtered over celite and washed with CH₂Cl₂. The filtrate was separated into its layers. The organic layer was dried (MgSO₄), filtered and the solvent was evaporated. The residue (6.7g) was purified by column chromatography over silica gel (eluent: CH₂Cl₂/CH₃OH/NH₄OH 95/5/0.2; 15-40 μm). The pure fractions were collected and the solvent was evaporated. This fraction was 10 crystallized from 2-propanone. The precipitate was filtered off and dried, yielding: 0.64g of compound 176.

Tables 4 to 13 list the compounds of formula (I) which were prepared according to one of the above examples.

15 Table 4

5

Comp No.	Ex. No.	a ·	Rá	R ^b	*	R ^C	Physical data
1	Bla	CH	Н	н	2.	н	HC1 (1:2)
2	Blb	СН	н	Н	2	**	
3	Blc	СН	н	н	2	CH2CH2NH2	HCl(1:4);H ₂ O(1:1)
4	Bla	СН	н	H	8	н	
5	Bla	СН	н	H	2	Н	
6	B5	СН	Н	Н	2	CH ₂ CH(2-propyl)NH ₂	
7	В3	СН	н	Н	8	CH(2-propyl)CH ₂ NH ₂	
8	В3	СН	н	Н	2	CH(2-propyl)CH ₂ NH ₂	H ₂ O (1:1)
9	Bla	СН	Н	8-Cl	2	Н .	HCl (1:2)
10	Blc	СН	н	н	8	CH ₂ CH(2-propyl)NH ₂	·
11	В3	СН	н	8-CI	2	CH(2-propyl)CH ₂ NH ₂	
12	Bla	СН	4-OH	Н	2	н	

Comp No.	Ex. No.	a	Rª	R ^b	*	R ^C	Physical data
13	В3	СН	н	8-C1	2	CH ₂ CH(2-propyl)NH ₂	
14	B6a	CH	3-OCH ₃	н	2	(C=O)OC(CH ₃) ₃	
15	Blc	СН	3-OCH ₃	н	2	CH ₂ CH(2-propyl)NH ₂	
16	B6a	N	3-CH ₃	н	2	***	
17	Bla	СН	H	н	8	н	HC1 (1:3)
18	Bla	N	Н	н	8	н	
19	Blc	N	Н	н	8	CH ₂ CH(2-propyl)NH ₂	HCl(1:3); H ₂ O(1:3)
20	Bla	N	3-OCH ₃	н	2	Н	
21	B4	N	3-OCH ₃	н	2	***	
22	Blc	N	3-OCH ₃	н	2	CH ₂ CH(2-propyl)NH ₂	
23	Bla	СН	н	н	4	Н	
24	B2	СН	н	н	4	CH₂CH₂NH₂	
88	Bla	N	2-CH ₃	3-CH ₃	8	н	
89	Blc	N	2-CH ₃	3-CH ₃	8	CH ₂ CH(2-propyl)NH ₂	HCl(1:4);H ₂ O(1:2)
90	Bla	CH	2-CH ₃	н	8	н	
91	Blc	СН	2-CH ₃	н	8	CH ₂ CH(2-propyl)NH ₂	H ₂ O (1:1)
92	B2	СН	2-CH ₃	н	8	CH₂CH₂NH₂	
104	B3	СН	н	н	8	CH ₂ CH(2-propyl)NH ₂	
105	В3	CH	н	н	8	CH(2-propyl)CH ₂ NH ₂	
106	B1c	N	3-CH ₃	н	2	CH ₂ CH(2-propyl)NH ₂	H ₂ 0 (1:2)
109	B5	СН	н	Н	8	***	
110	B5	N	2-CH ₃	3-CH ₃	8	***	
111	B5	СН	2-CH ₃	н	8	***	
112	B5	N	н	н	8	***	
113	B7	CH	н	Н	8	***	

- * position bicyclic heterocycle
- ** (CH₂)₂NH(C=O)OC(CH₃)₃
- *** CH₂CH(2-propyl)NH(C=O)OC(CH₃)₃

Table 5

			, R					
Comp No.	Ex. No.	a	Rª	R ^b	*	R ^c	G	Physical data
25	Bla	СН	Н	Н	2	Н	CHOC ₂ H ₅	
26	В3	СН	Н	н	2	CH(2-propyl)CH ₂ NH ₂	CHOC₂H₅	H ₂ O (1:1)
27	В3	СН	Н	Н	2	CH ₂ CH(2-propyl)NH ₂	CHOC₂H₅	
28	Bla	СН	Н	н	2	Н	***	
29	B3	СН	н	Н	2	CH(2-propyl)CH ₂ NH ₂	***	H ₂ O (1:1)
30	Bla	CH	н	Н	8	Н	***	·
31	В3	CH	н	Н	8	CH ₂ CH(2-propyl)NH ₂	***	
32	В3	СН	н	Н	8	CH(2-propyl)CH ₂ NH ₂	***	
33	Bla	СН	н	н	8	н	CHOC ₂ H ₅	
34	Bla	СН	3-OCH ₃	Н	2	н	CHOC ₂ H ₅	
35	Bla	N	н	н	2	н	CH₂	
36	B4	N	Н	н	2	**	CH₂	
37	Blc	N	н	н	2	CH ₂ CH(2-propyl)NH ₂	CH₂	HCl (1:4)
38	B4	CH	3-OCH₃	н	2	**	CHOC ₂ H ₅	
39 ⁽⁹⁾	Blc	СН	3-OCH ₃	н	2	CH ₂ CH(2-propyl)NH ₂	CHOC ₂ H ₅	HCl (1:3); H ₂ O (1:2)
40	B2	N	Н	Н	2	CH₂CH₂NH₂	CH ₂	
41	Bla	N	н	Ή.	8	Н	CH ₂	
42	Blc	N	Н	Н	8	CH ₂ CH(2-propyl)NH ₂	CH ₂	
43	Bla	CH	Н	CH ₃	8	Н	CH ₂	
44	Bla	CH	н	CH₃	8	Н	CHOC ₂ H ₅	
45	B2	N	Н	Н	8	CH ₂ CH ₂ NH ₂	CH ₂	
100	B3	СН	Н	Н	8	CH(2-propyl)CH ₂ NH ₂	CHOC ₂ H ₅	
107	Blc	CH	Н	H	8	CH ₂ CH(2-propyl)NH ₂	CHOC ₂ H ₅	
115	B5	CH	н	CH₃	8	CH(CH ₃) ₂ O C(CH ₃) ₃	CH₂	
116_	B1d	CH	Н	CH ₃	8	CH ₂ CH(2-propyl)NH ₂	CH ₂	

Comp	Ex.	a	Rª	R ^b	*	R ^c	G	Physical
No.	No.							data
117	Bld	СН	H	CH ₃	8	CH=O	CH ₂	
118	Bld	СН	H	CH ₃	8	CH2CH2NH2	***	H ₂ O(1:1)
119	Bld	СН	Н	CH ₃	8	CH ₂ CH(2-propyl)NH ₂	***	
120	В3	N	Н	CH ₃	8	CH2CH2NH2	CH ₂	HCl(1:4);
								H ₂ O(1:3)
121	B1d	СН	Н	CH ₃	8	CH=O	***	
122	Blc	N	н	CH ₃	8	CH ₂ CH(2-propyl)NH ₂	CH ₂	HCl(1:4);
			:					H ₂ O(1:1)
123	Bld	СН	н	CH ₃	8	CH₂CH₂NH₂	CH ₂	
124	Blc	СН	н	н	8	CH₂CH₂NH₂	***	HCl(1:3);
								H ₂ O(1:2)
125	Blc	СН	Н	CH ₃	8	CH₂CH₂NH₂	CHCH₃	H ₂ O(1:1)
126	Bld	СН	3-OCH₃	н	2	CH₂CH₂NH₂	CH₂	H ₂ O(1:2)
127	Blc	СН	4-CH ₃	H	2	CH ₂ CH(2-propyl)NH ₂	CH ₂	HCl(1:4);
}								H ₂ O(1:1)
128	Blc	CH	H	Н	8	CH₂CH₂NH₂	CH ₂	HCl(1:4);
								H ₂ O(1:1)
129	Blc	CH	Н	H	8	CH₂CH₂NH₂	CHCH ₃	H ₂ O(1:1)
130	B1c	CH	4-CH ₃	H	2	CH₂CH₂NH₂	CH ₂	HCl(1:4);
								H ₂ O(1:2)
131	Blc	CH	Н	Н	4	CH ₂ CH(2-propyl)NH ₂	CH₂	HCl(1:4);
								H ₂ O(1:2)
131	ВІЬ	CH	Н	CH ₃	8	C(CH ₃) ₃	CH ₂	
						N 0 (113)		
132	Blb	СН	H ·	Н	8	O Gray S	CH ₂	
						N C(CH ₃) ₃		
133	B2	CH	н	н	8	H	СНСН₃	HCl(1:2);
				7			,	H ₂ O(1:2)
134	Bic	СН	Н	н	2	CH₂CH₂NH₂	CHCH₃	H ₂ O(1:1)
135	В2	СН	4-CH ₃	H	2	Н	CH₂	HCl(1:2)
136	B2	N	Н	CH ₃	8	Н	CH ₂	,,
137	B11	СН	Н	Н	8	CH=O	CH ₂	

^{*} position quinoline

^{**} CH₂CH(2-propyl)NH(C=O)OC(CH₃)₃

^{***} CHO(CH₂)₂OC₂H₅

Table 6

Comp. No.	Ex. No.	*	G	Rª	Physical data
46	B5	2	CH ₂	CH ₂ CH(2-propyl)NH ₂	HCl(1:4);H ₂ O(1:1)
47	B5	8	CH ₂	CH ₂ CH(2-propyl)NH ₂	HCl(1:4);H ₂ O(1:1)
48	B5	8		Н	
49	B5	8	 	CH ₂ CH(2-propyl)NH ₂	H ₂ O (1:1)

^{*} position bicyclic heterocycle

5 <u>Table 7</u>

Co. No.	Ex. No.	*	a	Rª	G	R ^b	R°	Physical data
50	Bla	8	СН	н	-	Н	Н	
51	B5	8	СН	н	-	CH ₂ CH(2-propyl)NH ₂	Н -	
52	Bla	8	N	н	-	Н	Н	HCl (1:3)
53	B3	8	N	н	-	CH(2-propyl)CH ₂ NH ₂	Н	
54 ⁽³⁾	В3	8	N	н	-	CH ₂ CH(2-propyl)NH ₂	Н	H ₂ O (1:1)
55	Вбь	8	СН	2-Br	-,	**	Н	
56	Blc	8	СН	2-Вг	-	CH ₂ CH(2-propyl)NH ₂	Н	HCl(1:3);H₂O(1:3)
57	B6b	8	CH	2-CH₃		**	Н	

Co. No.	Ex. No.	*	a	Rª	G	R ^b	R°	Physical data
58	B1c	8	СН	2-CH ₃		CH ₂ CH(2-propyl)NH ₂	Н	HCl(1:4);H ₂ O(1:1)
59	Вба	2	СН	н	CH ₂	**	н	
60	Blc	2	СН	н	CH ₂	CH ₂ CH(2-propyl)NH ₂	Н	HCl(1:4);H ₂ O(1:1)
61	Вба	2	CH	з-осн	CH ₂	**	н	
62	Blc	2	CH	з-осн	CH ₂	CH ₂ CH(2-propyl)NH ₂	Н	HCl(1:4);H ₂ O(1:1)
63	B7	2	СН	з-он	CH ₂	CH ₂ CH(2-propyl)NH ₂	н	HCl(1:4);H ₂ O(1:1)
64	Bla	8	N	3-Cl	-	Н .	н	
65	B4	8	N	3-С1	-	**	н	
66	Blc	8	N	з-сі	-	CH ₂ CH(2-propyl)NH ₂	Н	HCl(1:3);H ₂ O(1:1)
67	B2	8	N	H	-	CH₂CH₂NH₂	Н	HCl(1:3);H ₂ O(1:3)
68	Bla	8	N	2-CI	-	Н	Н	
69	B4	8	N	2-CI	-	**	Н	
70 ⁽¹⁰⁾	Bic	8	N	2-Cl	-	CH ₂ CH(2-propyl)NH ₂	Н	HCl(1:3);H ₂ O(1:1)
139	B1c	5	N	з-сі	-	CH₂CH₂NH₂	CH ₃	HCl(1:3);H ₂ O(1:2)
140	Bld	5	N	н	-	CH2CH(2-propyl)NH2	CH ₃	
141	Blc	5	N	2-CI	-	CH₂CH₂NH₂	CH ₃	HCl(1:3);H ₂ O(1:3)
142	B1c	5	N	2-Cl		CH ₂ CH(2-propyl)NH ₂	CH ₃	•

- * position bicyclic heterocycle
- ** CH₂CH(2-propyl)NH(C=O)OC(CH₃)₃

Table 8

Ex. R^{b} $R^{\mathbf{a}}$ Comp. а G b R^c Physical data No No. 71 Н N Н N Н 72 S N Н Н HBr(1:2);H₂O(2:1) 73 Н Bla N Н

Comp. No	Ex. No.	a	ь	Rª	R ^b	G	R ^c	Physical data
74		N	N·	Н	Н	CH ₂	Н	
75		N	N	н	н	CH ₂	CH₂CH₂NH₂	H ₂ O (1:1)
76		O	СН	-	н	CH ₂	н	·
77		N ·	N	CH ₃	Н	CH ₂	н	
78	Blc	N	N	CH ₃	н	CH ₂	CH₂CH₂NH₂	
79		S	СН	-	н	CH₂	н	
80	Bla	s	N	-	н	CH ₂	н	HCl(1:2);H ₂ O(1:1)
81	B2	N	N	Н	H	-	CH₂CH₂NH₂	HCl(1:4)
82	Bla	N	N	H	OCH ₃	CH ₂	н	
83	В1ь	s	N	-	н	-	*	H ₂ O (1:1)
84	Blc	s	N	-	н	-	CH₂CH₂NH₂	HCl(1:3);H ₂ O(1:1)
85	Blb	N	N	CH₃	Н	CH₂	*	
86	В1ь	0	N	-	Н	-	*	
87	B1c	0	N		Н		CH ₂ CH ₂ NH ₂	

^{*} CH₂CH₂NH(C=O)OC(CH₃)₃

Table 9

Comp. No.	Ex. No.	Rª	Physical data
102	B1a	Н	HCl (1:3)
103	B5	CH ₂ CH(2-propyl)NH ₂	H ₂ O (1:1)

Table 10

Rnr

Comp.	Ex.	R ^b	R ^c	G-R ^a	Physical data
No.	No.				
93		н	Н	-CH ₂ -N	
101		CH₂CH₂NH₂	н	-CH ₂	
94		CH ₂ CH ₂ NH(C=O)O CH ₂ CH ₃	н	_CH ₂	
95		CH₂CH₂NH₂	н	-CH ₂	
96	Bla	H	н	-CH ₂ -CF ₃	
97	В2	CH₂CH₂NH₂	Н	$-H_2C$ N N CF_3	HCl(1:3);H₂O(1:1)
98	Bla	Н	Н	-CH ₂	
99	Blc	CH ₂ CH(2-propyl)NH ₂	Н	—сн ₂	HCl(1:3);H ₂ O(1:3)
108	В5	CH₂CH(2-propyl)NH₂	Н	-H ₂ C	

Rnr

Comp.	Ex.	R ^b	R°	G-Rª	Physical data
No.	No.				
114		*	Н	—CH ₂ —N—	
143	В6	CH ₂ CH(2-propyl)NH ₂	СН₃	-H ₂ C-	

^{*} $CH_2CH(2-propyl)NH(C=O)OC(CH_3)_3$

Table 11

5

Co No.	Ex. No.	a-a1-a2-a3	*	Rª	R°	R ^b	G	Physical data
144	B1c	CH=N-CH=C	8	Н	-	CH ₂ CH(2-propyl)NH ₂	CH₂	HCl(1:3);
								H ₂ O(1:4)
145	Blc	CH=C-N=C	8	Н	Н	CH ₂ CH(2-propyl)NH ₂	CH₂	HCl(1:3);
								H ₂ O(1:2)
146	Blc	CH=C-C=N	8	-	Н	CH ₂ CH(2-propyl)NH ₂	CH ₂	HCl(1:3);
	Ì	•						H ₂ O(1:2)
147	B2	CH=C-CH=C	8	CH₃	Cl	Н	CH₂	
148	B3	CH=N-CH=C	8	н	-	CH₂CH₂NH₂	CHOC₂H₅	
149	B2	CH=C-CH=N	8	-	Н	н	CH ₂	HCl(1:2);
Ì								H ₂ O(1:1)
150	Blc	CH=C-CH=C	7	CH₃	Cl	CH ₂ CH ₂ NH ₂	CH ₂	HCl(1:4);
		,						H ₂ O(1:2)
151	В3	CH=N-CH=C	8	н		CH₂CH₂NH₂	CH₂	

Co No.	Ex. No.	a-a1-a2-a3	*	Rª	R°	R ^b	G	Physical data
152	B2	CH=N-CH=C	8	Н	-	Н	CH₂	HCl(1:4);
								H ₂ O(1:2)
153	B3	CH=C-CH=N	8	-	H	CH ₂ CH ₂ NH ₂	CHOC2H5	

• position bicyclic heterocycle

Co ¹ No.	Ex. No.	Rª	R ^b	Physical data
154	Blc	Н	3-propylamine	HCl(1:3);H ₂ O(1:1)
155	Віь	Н	(H ₃ C) ₃ C O H NH ₂	
156	Ble	н	H ₂ N NH	
157	В7с	Н		trans
			(H ₃ C) ₃ C HN	
158	В7с	Н	H ₃ C NH	
159	B9a	н	2-ethylamipe	
160	Blc	Н	3-propylmethylamine	
161	Blc	Н	H ₂ N N	cis;HCl(1:3);H ₂ O(1:1)
162	Blc	Н	CH ₃	HCl(1:4);H₂O(1:1)
163	B4	н	3-isobutylamine	
164	Blc	н	2-ethylmethylamine	HCI(1:2)

Co	Ex.	Rª	R ^b	Physical
No.	No.	K	K.	data
165	Bla	Н	\bigcirc	trans;H ₂ O(1:1)
166	DO:	CH ₃	H ₂ N	
166	B9a	Cn ₃	2-ethylamine	
167	В9ь	Н	H ₂ N	cis
168	В7с	н		cis
169	В3	H	H ₂ N CH ₂	HCl(1:3);H₂O(1:2)
170	B10	Н	H_{2N} H_{2N} CH_{2}	
171	B10	Н	H ₂ N CH CH ₃) ₂	H ₂ O(1:1)
172	Blc	н	H_2N N CH_2	HCl(1:4);H ₂ O(1:2)
173	В9с	Н	HN CH ₂ —	HCl(1:3)H ₂ O(1:2)
174	Blc	Н	H ₂ N CH CH ₃	
175	В7с	н		cis
176	B12	н	(H ₃ C) ₃ C—O—HN	
			CH ₃	

Table 13

Co No.	Ex. No.	G	L	a.	R _a .	Physical data
177	Bld	2-ethylamine	N CH ₃	CH	Н	HCl(1:3);H ₂ O (1:3)
178	Blc	2-ethylamine	CH ₃	N	Н	HCl(1:4);H₂O (1:4)
179	Blc	2-ethylamine	N CH ₃	СН	СН3	H ₂ O(1:1)
180	В1ь	(H ₃ C) ₃ C O H ₂ CH ₂	N CH ₃	СН	Н	
181	Blc	CH(CH ₃) ₂ H ₂ N CH ₂ —		СН	Н	HCl(1:3);H₂O (1:2)
182	Blc	2-ethylamine	NH O	СН	Н	HCl(1:3);H₂O (1:2)
183	Bic	2-ethylamine		СН	н	
184	Blc	2-ethylamine		СН	Н	HCl(1:4);H₂O (1:1)
185	Bld	2-ethylamine		СН	Н	C ₂ H ₂ O ₄ (2:7)

Table 14: Physical data

Comp.	(·		<u> </u>		N	melting point
No.					İ		
	Theor.	Ехр.	Theor.	Exp.	Theor.	Exp.	
1	61.40	60.70	5.85	6.04	16.27	15.54	
3	51.08	51.16	6.07	6.35	14.89	14.17	
4	73.92	73.29	6.49	6.52	19.59	19.38	206°C
6	73.27	73.12	7.74	7.73	18.99	18.77	135°C
7	73.27	71.85	7.74	7.80	18.99	18.61	188°C
8	70.40	69.73	7.88	7.40	18.24	17.56	80°C
9							> 250°C
10	73.27	72.82	7.74	7.58	18.99	18.63	172°C
11							190°C
13	67.98	66.43	6.97	6.79	17.62	17.02	164°C
15	71.16	70.66	7.68	7.58	17.78	17.81	210°C
19	51.45	51.64	6.97	6.89	16.15	15.96	240°C
22	68.47	68.04	7.45	7.52	20.70	20.55	206°C
23	73.92	71.70	6.49	6.53	19.59	19.92	140°C
24	71.97	69.89	7.05	7.10	20.98	20.07 .	196°C
89	51.46	53.22	6.94	7.11	15.00	15.14	24°C
91	70.85	69.82	8.07	8.29	17.71	17.48	180°C
92	72.43	71.51	7.29	7.30	20.27	20.10	176°C
104	72.87	70.26	7.53	7.27	19.61	18.73	88°C
105	72.87	71.37	7.53	7.39	19.61	19.39	135°C
106	65.69	66.19	7.96	7.62	19.86	19.71	110°C
26	69.02	69.16	7.99	7.68	16.65	16.79	140°C
27	71.57	70.60	7.87	7.80	17.27	17.14	166°C
29	67.86	67.64	8.08	7.79	15.32	15.15	100°C
31	70.16	68.97	7.98	7.97	15.84	15.56	110°C
32	70.16	69.35	7.98	8.34	15.84	14.73	98°C
33	71.79	70.72	6.78	7.17	17.44	16.69	145°C
37							215°C
39							209°C
40	68.80	66.01	6.78	6.60	24.42	23.31	138°C
42	70.40	69.14	7.50	7.50	22.10	21.68	180°C
43	74.36	73.02	6.78	6.65	18.85	18.41	155°C
44	72.26	71.53	7.03	7.26	16.85	16.40	186°C
45	68.80 ·	66.74	6.78	6.64	24.42	23.77	178°C
100	71.57	71.16	7.87	7.93	17.27	17.44	202°C
107	71.57	69.77	7.87	7.85	17.27	16.40	78°C

Comp.	С		Н		N		melting poin
No.	Theor.	Eva	Theor.	Exp.	Theor.	Exp.	
46	THEOL.	Exp.	Theor.	Ехр.	THEOI.	Exp.	230°C
47							1
	72.50	71 54	7.05	7.12	20.16	10.01	230°C
48	72.59	71.54	7.25	7.13	20.16	19.91	205°C
49	69.30	70.08	8.50	8.37	18.65	18.93	140°C
51	72.19	70.66	8.39	8.43	19.43	18.79	120°C
53	69.25	68.88	8.14	8.28	22.61	22.23	
54	66.49	66.30	8.26	7.77	21.71	21.53	144°C
56	46.27	47.19	6.57	6.44	12.45	12.16	> 250°C
58				ı			210°C
60							212°C
62	52.51	53.38	7.24	7.63	13.12	12.37	240°C
63	51.76	52.74	7.08	7.32	13.41	12.93	240°C
66	50.43	50.60	6.60	6.58	16.47	16.28	> 250°C
67	47.62	46.73	6.90	6.83	17.67	17.19	230°C
70							238°C
80							210°C
81	48.38	47.77	5.61	5.61			1
82	67.00	66.51	6.43	6.29	22.32	22.12	1
83	61.15	62.11	6.71	6.60	16.46	16.88	
84	48.51	48.46	5.62	5.35	16.16	16.03	
87	67.00	66.42	6.43	6.55	22.32	21.80	
103	68.78	68.77	8.31	8.23	19.25	18.78	88°C
96	58.73	58.59	5.16	5.03	22.83	22.40	144°C
97							210°C
99	53.51	52.63	7.15	7.02	13.87	13.24	200°C
108	70.08	68.99	7.92	8.10	22.00	21.65	160°C
116							203°C
117							218°C
141		:					225°C
177							>260°C
139					1		190°C
118							48°C
144							220°C
143	70.55	66.03	8.11	8.14	21.33	18.98	
119							145°C
121							185°C
140						•	172°C
120							210°C

Comp.	С]	Н		N	melting point
No.				· · · · · · · · · · · · · · · · · · ·			
ļ	Theor.	Exp.	Theor.	Exp.	Theor.	Exp.	·
142							98°C
122							245°C
154							90°C
145							190°C
123							194°C
124							150°C
146							240°C
125							74°C
178				·			160°C
150							>250°C
126							90°C
127							200°C
128			·				210°C
157							185°C
159							140°C
151							212°C
160	73.02	72.95	6.71	6.70	20.27	20.35	
129							170°C
130							150°C
131							>250°C
152							230°C
153							169°C
131							120°C
161							206°C
132							160°C
133							210°C
134							81°C
162							210°C
147							>250°C
163							168°C
179							116°C
135	62.16	62.10	6.12	6.06	15.76	15.71	
164							146°C
136			i			,	188°C
165							112°C
166							114°C
149							210°C
180							247°C

Comp. No.	C		Н		N		melting point
	Theor.	Exp.	Theor.	Ехр.	Theor.	Ехр.	
167							167°C
181							235°C
182							>250°C
184	47.75	47.58	6.01	6.37	17.72	17.00	
169							180°C
170							73°C
171							72°C
172							178°C
173							190°C
137							196°C
175							228°C
176							168°C
185			<u> </u>				158°C

C. Pharmacological example

Example C1: In vitro screening for activity against Respiratory Syncytial Virus.

The percent protection against cytopathology caused by viruses (antiviral activity or IC₅₀) achieved by tested compounds and their cytotoxicity (CC₅₀) were both calculated from dose-response curves. The selectivity of the antiviral effect is represented by the selectivity index (SI), calculated by dividing the CC₅₀ (cytotoxic dose for 50% of the cells) by the IC₅₀ (antiviral activity for 50 % of the cells).

10 Automated tetrazolium-based colorimetric assays were used for determination of IC50 and CC50 of test compounds. Flat-bottom, 96-well plastic microtiter trays were filled with 180 µl of Eagle's Basal Medium, supplemented with 5 % FCS (0% for FLU) and 20 mM Hepes buffer. Subsequently, stock solutions (7.8 x final test concentration) of compounds were added in 45 µl volumes to a series of triplicate wells so as to allow 15 simultaneous evaluation of their effects on virus- and mock-infected cells. Five fivefold dilutions were made directly in the microtiter trays using a robot system. Untreated virus controls, and HeLa cell controls were included in each test. Approximately 100 TCID₅₀ of Respiratory Syncytial Virus was added to two of the three rows in a volume of 50 µl. The same volume of medium was added to the third row to measure the 20 cytotoxicity of the compounds at the same concentrations as those used to measure the antiviral activity. After two hours of incubation, a suspension (4 x 10⁵ cells/ml) of HeLa cells was added to all wells in a volume of 50µl. The cultures were incubated at

10

37°C in a 5% CO₂ atmosphere. Seven days after infection the cytotoxicity and the antiviral activity was examined spectrophotometrically. To each well of the microtiter tray, 25 μl of a solution of MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) was added. The trays were further incubated at 37°C for 2 hours, after which the medium was removed from each cup. Solubilization of the formazan crystals was achieved by adding 100 μl 2-propanol. Complete dissolution of the formazan crystals were obtained after the trays have been placed on a plate shaker for 10 min. Finally, the absorbances were read in an eight-channel computer-controlled photometer (Multiskan MCC, Flow Laboratories) at two wavelengths (540 and 690 nm). The absorbance measured at 690 nm was automatically subtracted from the absorbance at 540 nm, so as to eliminate the effects of non-specific absorption.

Particular IC₅₀, CC₅₀ and SI values are listed in Table 15 hereinbelow. Table 15

Co. No.	IC ₅₀ (μM)	CC ₅₀ (µM)	SI
42	0.0004	>10.05	>25119
31	0.0008	12.68	15849
56	0.0016	12.71	7943
145	0.00631	25.12	3981
6	0.0126	10.00	794
156	0.01259	19.95	1585
131	0.0316	19.94	631
53	0.1259	>9.95	>79
29	0.3162	10.12	32
148	1	25	25
97	1.5849	>99.85	>63

Claims

10

15

25

1. A compound of formula

$$Q = \begin{pmatrix} R^1 & & & \\ & & & \\ Q & & & \\ & & & \\ Q & & & \\ & & &$$

a prodrug, N-oxide, addition salt, quaternary amine, metal complex or stereochemically

5 isomeric form thereof wherein

-a¹=a²-a³=a⁴- represents a bivalent radical of formula

-CH=CH-CH=CH- (a-1); -N=CH-CH=CH- (a-2); -CH=N-CH=CH- (a-3); -CH=CH-N=CH- (a-4); or -CH=CH-CH=N- (a-5);

wherein each hydrogen atom in the radicals (a-1), (a-2), (a-3), (a-4) and (a-5) may optionally be replaced by halo, C₁₋₆alkyl, nitro, amino, hydroxy, C₁₋₆alkyloxy, polyhaloC₁₋₆alkyl, carboxyl, aminoC₁₋₆alkyl, mono- or

 $di(C_{1-4}alkyl)aminoC_{1-6}alkyl, C_{1-6}alkyloxycarbonyl, hydroxyC_{1-6}alkyl, or a radical of formula$

wherein =Z is =O, =CH-C(=O)-NR 5a R 5b , =CH₂, =CH-C₁₋₆alkyl, =N-OH or =N-O-C₁₋₆alkyl;

20 Q is a radical of formula

wherein Alk is C₁₋₆alkanediyl;

Y¹ is a bivalent radical of formula $-NR^2$ - or $-CH(NR^2R^4)$ -; X¹ is NR^4 , S, S(=O), S(=O)₂, O, CH₂, C(=O), C(=CH₂), CH(OH), CH(CH₃), CH(OCH₃), CH(SCH₃), CH(NR^{5a}R^{5b}), CH₂-NR⁴ or NR⁴-CH₂:

 X^2 is a direct bond, CH_2 , C(=O), NR^4 , C_{1-4} alkyl- NR^4 , NR^4 - C_{1-4} alkyl; t is 2, 3, 4 or 5; u is 1, 2, 3, 4 or 5; v is 2 or 3; and

whereby each hydrogen atom in Alk and the carbocycles and the heterocycles defined in radicals (b-3), (b-4), (b-5), (b-6), (b-7) and (b-8) may optionally be replaced by R³; with the proviso that when R³ is hydroxy or C₁₋₆alkyloxy, then R³ can not replace a hydrogen atom in the α position relative to a nitrogen atom;

G is a direct bond or C_{1-10} alkanediyl optionally substituted with one, two or three substituents selected from hydroxy, C_{1-6} alkyloxy, aryl C_{1-6} alkyloxy, C_{1-6} alkylthio, aryl C_{1-6} alkylthio, arylcarbonyl, HO(-CH₂-CH₂-O)_n-, C_{1-6} alkyloxy(-CH₂-CH₂-O)_n-, arryl C_{1-6} alkyloxy(-CH₂-CH₂-O)_n-, amino, mono-or di(C_{1-6} alkyloxycarbonylamino and aryl;

R¹ is a bicyclic heterocycle selected from quinolinyl, quinoxalinyl, benzofuranyl,
benzothienyl, benzimidazolyl, benzoxazolyl, benzthiazolyl, pyridopyridyl,
naphthiridinyl, 1*H*-imidazo[4,5-b]pyridinyl, 3*H*-imidazo[4,5-b]pyridinyl,
imidazo[1,2-a]pyridinyl, 2,3-dihydro-1,4-dioxino[2,3-b]pyridyl or a radical of formula

$$(CH_{2})_{m} \qquad (CH_{2})_{m} \qquad (CH_{2})_{p} \qquad (CH_$$

and said bicyclic heterocycles may optionally be substituted in either of the two cycles
with 1 or where possible more, such as 2, 3 or 4, substituents selected from halo,
hydroxy, amino, cyano, carboxy, C₁₋₆alkyl, C₁₋₆alkyloxy, C₁₋₆alkylthio, C₁₋₆alkyloxyC₁₋₆alkyl, aryl, arylC₁₋₆alkyl, arylC₁₋₆alkyloxy, hydroxyC₁₋₆alkyl, mono-or di(C₁₋₆alkyl)amino, mono-or di(C₁₋₆alkyl)aminoC₁₋₆alkyl, polyhaloC₁₋₆alkyl, C₁₋₆alkylcarbonylamino,
C₁₋₆alkyl-SO₂-NR^{5c}-, aryl-SO₂-NR^{5c}-, C₁₋₆alkyloxycarbonyl, -C(=O)-NR^{5c}R^{5d}, HO(CH₂-CH₂-O)_n-, halo(-CH₂-CH₂-O)_n-, C₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, arylC₁₋₆alkyloxy(CH₂-CH₂-O)_n- and mono-or di(C₁₋₆alkyl)amino(-CH₂-CH₂-O)_n-;
each n independently is 1, 2, 3 or 4;
each m independently is 1 or 2;

each p independently is 1 or 2;

each R^2 independently is hydrogen, formyl, $C_{1\text{-}6}$ alkylcarbonyl, Hetcarbonyl, pyrrolidinyl, piperidinyl, homopiperidinyl, $C_{3\text{-}7}$ cycloalkyl substituted with $N(R^6)_2$, or $C_{1\text{-}10}$ alkyl substituted with $N(R^6)_2$ and optionally with a second, third or fourth

- substituent selected from amino, hydroxy, C₃₋₇cycloalkyl, C₂₋₅alkanediyl, piperidinyl, mono-or di(C₁₋₆alkyl)amino, C₁₋₆alkyloxycarbonylamino, aryl and aryloxy; R³ is hydrogen, hydroxy, C₁₋₆alkyl, C₁₋₆alkyloxy, arylC₁₋₆alkyl or arylC₁₋₆alkyloxy; R⁴ is hydrogen, C₁₋₆alkyl or arylC₁₋₆alkyl;
 - $R^{5a},\,R^{5b},\,R^{5c}$ and R^{5d} each independently are hydrogen or $C_{1\text{-}6}alkyl;$ or
- 10 R^{5a} and R^{5b}, or R^{5c} and R^{5d} taken together form a bivalent radical of formula -(CH₂)_s-wherein s is 4 or 5;
 - R^6 is hydrogen, C_{1-6} alkyl, formyl, hydroxy C_{1-6} alkyl, C_{1-6} alkylcarbonyl or C_{1-6} alkyloxycarbonyl;
- aryl is phenyl or phenyl substituted with 1 or more, such as 2, 3 or 4, substituents selected from halo, hydroxy, C₁₋₆alkyl, hydroxyC₁₋₆alkyl, polyhaloC₁₋₆alkyl, and C₁₋₆alkyloxy;

Het is pyridyl, pyrimidinyl, pyrazinyl, pyridazinyl.

- A compound according to claim 1 wherein -a¹=a²-a³=a⁴- is a radical of formula
 (a-1), (a-2) or (a-3).
 - 3. A compound according to claim 1 or 2 wherein Q is a radical of formula (b-5) wherein v is 2 and Y¹ is -NR²-.
- 25 4. A compound according to anyone of claims 1 to 3 wherein R² is C₁₋₁₀alkyl substituted with NHR⁶.
- A compound according to anyone of claims 1 to 4 wherein G is a direct bond or C₁₋₁₀alkanediyl optionally substituted with one, two or three substituents selected from hydroxy, C₁₋₆alkyloxy, arylC₁₋₆alkyloxy, HO(-CH₂-CH₂-O)_n-, C₁₋₆alkyloxy(-CH₂-CH₂-O)_n-, arylC₁₋₆alkyloxy(-CH₂-CH₂-O)_n-.
 - 6. A compound according to claim 1 wherein the compound is selected from (±)-N-[1-(2-aminoethyl)-4-piperidinyl]-4-methyl-1-[1-(8-quinolinyl)ethyl]-1H-
- benzimidazol-2-amine monohydrate; $(\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(2-bromo-5,6,7,8-tetrahydro-8-quinolinyl)-<math>IH$ -benzimidazol-2-amine trihydrochloride trihydrate; $(\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-[(2-ethoxyethoxy)-8-quinolinylmethyl]-4-methyl-<math>IH$ -benzimidazol-

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2-amine; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(2-chloro-5,6,7,8tetrahydro-5-quinoxalinyl)-1H-benzimidazol-2-amine trihydrochloride trihydrate; (\pm) -N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-[(1-methyl-1H-benzimidazol-4yl)methyl]-1H-benzimidazol-2-amine; (±)-N-[1-(2-amino-3-methylbutyl)-4-5 piperidinyl]-1-(ethoxy-8-quinolinylmethyl)-1H-benzimidazol-2-amine; (\pm)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-4-methyl-1-(5,6,7,8-tetrahydro-5quinoxalinyl)-1H-benzimidazol-2-amine; (±)-N-[1-(2-aminoethyl)-4-piperidinyl]-7-methyl-3-(8-quinolinylmethyl)-3H-imidazo[4,5-b]pyridin-2-amine tetrahydrochloride trihydrate; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-7-10 methyl-3-(8-quinolinylmethyl)-3H-imidazo[4,5-b]pyridin-2-amine tetrahydrochloride monohydrate; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(8-quinolinylmethyl)-1H-imidazo[4,5-c]pyridin-2-amine trihydrochloride dihydrate; N-[1-(2-aminoethyl)-4-piperidinyl]-4-methyl-1-(8-quinolinylmethyl)-1H-benzimidazol-2-amine; N-[1-(8-quinolinylmethyl)-1H-benzimidazol-2-yl]-1,3-15 propanediamine trihydrochloride monohydrate; (±)-N-[1-(2-aminoethyl)-4piperidinyl]-1-[(2-ethoxyethoxy)-8-quinolinylmethyl]-1H-benzimidazol-2-amine trihydrochloride dihydrate; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(8quinolinylmethyl)-1H-imidazo[4,5-b]pyridine-2-amine trihydrochloride dihydrate; (±)-N-[1-[1-(aminomethyl)-2-methylpropyl]-4-piperidinyl]-1-[(2-ethoxyethoxy)-8-20 quinolinylmethyl]-1H-benzimidazol-2-amine; (±)-N-[1-(2-aminoethyl)-4piperidinyl]-3-(2-quinolinylmethyl)-3H-imidazo[4,5-b]pyridin-2-amine trihydrochloride trihydrate; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(1isoquinolinylmethyl)-1H-benzimidazol-2-amine trihydrochloride trihydrate; N-[1-(2-aminoethyl)-4-piperidinyl]-1-(5,6,7,8-tetrahydro-5-quinoxalinyl)-1H-25 benzimidazol-2-amine trihydrochloride trihydrate; (±)-N-[1-(2-amino-3methylbutyl)-4-piperidinyl]-3-(quinolinylmethyl)-3H-imidazo[4,5-b]pyridin-2amine; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-4-methyl-1-(8quinolinylmethyl)-1H-benzimidazol-2-amine; (±)-N-[1-(2-aminoethyl)-4piperidinyl]-1-(2-chloro-5,6,7,8-tetrahydro-5-quinoxalinyl)-4-methyl-1H-30 benzimidazol-2-amine trihydrochloride.trihydrate; (±)-N-[1-(2-aminoethyl)-4piperidinyl]-1-(5,6,7,8-tetrahydro-2,3-dimethyl-5-quinoxalinyl)-1H-benzimidazol-2-amine trihydrochloride trihydrate; (±)-N-[1-(2-amino-3-methylbutyl)-4piperidinyl]-1-[(2-ethoxyethoxy)-8-quinolinylmethyl]-1H-benzimidazol-2-amine; (\pm) -N-[1-(2-amino-3-methylbutyl)-4-piperidinyl-1-(3-chloro-5,6,7,8-tetrahydro-5-35 quinoxalinyl)-1H-benzimidazol-2-amine trihydrochloride monohydrate; (±)-N-[1-(2-aminoethyl)-4-piperidinyl]-1-(3-chloro-5,6,7,8-tetrahydro-5-quinoxalinyl)-4methyl-1H-benzimidazol-2-amine trihydrochloride dihydrate; (±)-N-[1-(2-

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aminoethyl)-4-piperidinyl]-1-[(2-ethoxyethoxy)-8-quinolinylmethyl]-4-methyl-1H-benzimidazol-2-amine monohydrate; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-3-(8-quinolinylmethyl)-3H-imidazo[4,5-c]pyridin-2-amine trihydrochloride tetrahydrate; (±)-N-[1-(2-aminoethyl)-4-piperidinyl]-3-(8-quinolinylmethyl)-3H-imidazo[4,5-b]pyridin-2-amine; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-4-methyl-1-[(1-methyl-1H-benzimidazol-4-yl)methyl]-1H-benzimidazol-2-amine; (±)-N-[1-(2-amino-3-methylbutyl)-4-piperidinyl]-1-(2-chloro-5,6,7,8-tetrahydro-5-quinoxalinyl)-4-methyl-1H-benzimidazol-2-amine; a prodrug, N-oxide, addition salt, quaternary amine, metal complex or stereochemically isomeric form thereof.

- 7. A compound according to any one of claims 1 to 6 for use as a medicine.
- 8. A pharmaceutical composition comprising a pharmaceutically acceptable carrier, and as active ingredient a therapeutically effective amount of a compound as described in any one of claims 1 to 6.
 - 9. A process of preparing a composition as claimed in claim 8, <u>characterized in that</u>, a pharmaceutically acceptable carrier is intimately mixed with a therapeutically effective amount of a compound as described in any one of claims 1 to 6.
 - 10. An intermediate of formula

$$P = Q_1 = N = A \begin{pmatrix} A & A \\ A & A \end{pmatrix} \begin{pmatrix} A & A \\ A \end{pmatrix} \begin{pmatrix} A & A \\ A \end{pmatrix} \begin{pmatrix} A & A \\ A \end{pmatrix} \begin{pmatrix} A &$$

with R^1 , G and $-a^1=a^2-a^3=a^4$ defined as in claim 1, P being a protective group, and Q_1 being defined as Q according to claim 1 but being devoided of the R^2 or R^6 substituent.

11. An intermediate of formula

with R^1 , G and $-a^1=a^2-a^3=a^4$ defined as in claim 1, and $(O=)Q_3$ being a carbonyl derivative of Q, said Q being defined according to claim 1, provided that it is devoided of the NR^2R^4 or NR^2 substituent.

5 12. An intermediate of formula

$$Q = N$$

$$Q = N$$

$$Q = N$$

$$Q = A$$

$$Q$$

with R^1 , Q and $-a^1=a^2-a^3=a^4$ defined as in claim 1, and $(O=)G_2$ being a carbonyl derivative of G, said G being defined according to claim 1.

10 13. A process of preparing a compound as claimed in claim 1, characterized by,

a) reacting an intermediate of formula (II-a) or (II-b) with an intermediate of formula (III)

with R¹, G, Q and -a¹=a²-a³=a⁴- defined as in claim 1, and W₁ being a suitable leaving group, in the presence of a suitable base and in a suitable reaction-inert solvent;

b) deprotecting an intermediate of formula (IV)

$$P = Q_1 = \begin{bmatrix} Q & R^1 & Q_1 &$$

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with R^1 , G, and $-a^1=a^2-a^3=a^4$ - defined as in claim 1, H-Q₁ being defined as Q according to claim 1 provided that R^2 or at least one R^6 substituent is hydrogen, and P being a protective group;

5 c) deprotecting and reducing an intermediate of formula (IV-a)

with R^1 , G, and $-a^1=a^2-a^3=a^4$ - defined as in claim 1, H-Q₁ being defined as Q according to claim 1 provided that R^2 or at least one R^6 substituent is hydrogen, Q_{1a}(CH=CH) being defined as Q₁ provided that Q₁ comprises an unsaturated bond, and P being a protective group;

d) deprotecting an intermediate of formula (V)

with R^1 , G, and $-a^1=a^2-a^3=a^4$ - defined as in claim 1, and H_2N-Q_2 being defined as Q according to claim 1 provided that both R^6 substituents are hydrogen or R^2 and R^4 are both hydrogen;

e) deprotecting an intermediate of formula (VI)

$$P = Q_{2} = \begin{pmatrix} R^{1} & & & \\$$

with R¹, G, and -a¹=a²-a³=a⁴- defined as in claim 1, and H₂N-Q₂ being defined as Q according to claim 1 provided that both R⁶ substituents are hydrogen or R² and R⁴ are both hydrogen, and P being a protective group;

f) deprotecting an intermediate of formula (VII) or (VIII)

$$P = Q_{1} \cdot (OP) \longrightarrow \begin{pmatrix} R^{1} \\ A^{2} \\ A^{3} \end{pmatrix} \longrightarrow H = Q_{1} \cdot (OH) \longrightarrow \begin{pmatrix} R^{1} \\ A^{2} \\ A^{3} \end{pmatrix}$$

$$(VIII) \qquad (I-a-2)$$

$$Q^{R^{1}} \longrightarrow H_{2}N = Q_{2} \cdot (OH) \longrightarrow \begin{pmatrix} R^{1} \\ A^{2} \\ A^{3} \end{pmatrix}$$

$$(VIII) \qquad (I-a-1-1)$$

with R^1 , G, and $-a^1=a^2-a^3=a^4$ - defined as in claim 1, H-Q₁·(OH) being defined as Q according to claim 1 provided that R^2 or at least one R^6 substituent is hydrogen and provided that Q comprises a hydroxy moiety, H₂N-Q₂·(OH) being defined as Q according to claim 1 provided that both R^6 substituents are hydrogen or R^2 and R^4 are both hydrogen and provided that Q comprises a hydroxy moiety, and P being a protective group;

g) amination of an intermediate of formula (IX)

$$Q = Q_3$$
 Q_3
 $Q = Q_3$
 $Q = Q_$

10

15

5

with R^1 , G, and $-a^1=a^2-a^3=a^4$ - defined as in claim 1, and H_2N-Q_3H being defined as Q according to claim 1 provided that both R^6 substituents are hydrogen or R^2 and R^4 are both hydrogen, and the carbon adjacent to the nitrogen carrying the R^6 , or R^2 and R^4 substituents contains at least one hydrogen, in the presence of a suitable amination reagent;

h) reducing an intermediate of formula (X)

NC-Q₄

$$R^1$$
reduction
 H_2N -CH₂-Q₄
 R^1
 A^1
 A^2
 A^2
 A^3
(I-a-1-3)

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with R^1 , G, and $-a^1=a^2-a^3=a^4$ defined as in claim 1, and $H_2N-CH_2-Q_4$ being defined as Q according to claim 1 provided that Q comprises a $-CH_2-NH_2$ moiety, in the presence of a suitable reducing agent;

i) reducing an intermediate of formula (X-a)

with G, and $-a^1=a^2-a^3=a^4$ - defined as in claim 1, $H_2N-CH_2-Q_4$ being defined as Q according to claim 1 provided that Q comprises a $-CH_2-NH_2$ moiety, and R^1 being defined as R^1 according to claim 1 provided that it comprises at least one substituent, in the presence of a suitable reducing agent and suitable solvent;

10 j) amination of an intermediate of formula (XI)

with R^1 , G, and $-a^1=a^2-a^3=a^4$ defined as in claim 1, and H_2N -CH₂-CHOH-CH₂-Q₄ being defined as Q according to claim 1 provided that Q comprises a CH₂-CHOH-CH₂-NH₂ moiety, in the presence of a suitable amination reagent;

k) reacting an intermediate of formula (XII) with formic acid, formamide and ammonia

$$C_{1^{-4}}\text{alkyl} = C_{1^{-4}}\text{alkyl} = C_{1^{-$$

with R¹, G, and -a¹=a²-a³=a⁴- defined as in claim 1, and H-C(=O)-Q₁ being defined as Q according to claim 1 provided that R² or at least one R⁶ substituent is formyl;

1) amination of an intermediate of formula (XIII) by reaction with an intermediate of formula (XIV)

$$(O=)Q_{5} \xrightarrow{R^{1}} A_{3}^{1} + R^{2a} \xrightarrow{NH_{2}} A_{3}^{2a} + R^{2a} \xrightarrow{NH_{2}} A_{3}^{2a}$$

$$(XIII) \qquad (XIV) \qquad (I-c)$$

with R^1 , G, and $-a^1=a^2-a^3=a^4$ defined as in claim 1, and R^{2a} -NH-HQ₅ being defined as Q according to claim 1 provided that R^2 is other than hydrogen and is represented by R^{2a} , R^4 is hydrogen, and the carbon atom adjacent to the nitrogen atom carrying the R^2 and R^4 substituents, carries also at least one hydrogen atom, in the presence of a suitable reducing agent;

m) reducing an intermediate of formula (XV)

with R¹, G, and -a¹=a²-a³=a⁴- defined as in claim 1, and

(R⁶)₂N-[(C₁₋₉alkyl)CH₂OH]-NH-HQ₅ being defined as Q according to claim 1 provided that R² is other than hydrogen and is represented by C₁₋₁₀alkyl substituted with N(R₆)₂ and with hydroxy, and the carbon atom carrying the hydroxy, carries also two hydrogen atoms, and provided that R⁴ is hydrogen, and the carbon atom adjacent to the nitrogen atom carrying the R² and R⁴ substituents, carries also at least one hydrogen atom, with a suitable reducing agent;

n) deprotecting an intermediate of formula (XVI), (XVI-a) or (XVI-b)

$$P = Q_{1} = \begin{pmatrix} A & O & P \end{pmatrix}_{w}$$

$$Q_{1} = \begin{pmatrix} A & O & H \end{pmatrix}_{w}$$

$$Q_{1} = \begin{pmatrix} A & O & H \end{pmatrix}_{w}$$

$$Q_{1} = \begin{pmatrix} A & O & H \end{pmatrix}_{w}$$

$$A & A & A & A \\
A & A &$$

10

15

$$\begin{array}{c} P_1 - O \\ O \\ A \\ R_{1a'} \end{array}$$

$$\begin{array}{c} P - Q_1 \\ O \\ A \\ R_{1a'} \end{array}$$

$$\begin{array}{c} A \\ A \\ R_{1a'} \end{array}$$

$$\begin{array}{c} A \\ A \\ R_{1a'} \end{array}$$

$$\begin{array}{c} (I-d-1) \\ O \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} O \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} A \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} O \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} O \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} O \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} O \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} O \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} O \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} O \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \\ A \\ R_{1a'-A} - O - H \end{array}$$

$$\begin{array}{c} O \\ A \\ R_{1a'-A} - O - H \\ R_{$$

with G, and $-a^1=a^2-a^3=a^4$ defined as in claim 1, and H-Q₁ being defined as Q according to claim 1 provided that R² or at least one R⁶ substituent is hydrogen, and R^{1a}-(A-O-H)_w, R^{1a'}-(A-O-H)₂ and R^{1a''}-(A-O-H)₃ being defined as R¹ according to claim 1 provided that R¹ is substituted with hydroxy, hydroxyC₁₋₆alkyl, or HO(-CH₂-CH₂-O)_n-, with w being an integer from 1 to 4 and P or P₁ being a suitable protecting group, with a suitable acid.

o) amination of an intermediate of formula (XVII)

$$C_{1^{-4}alkyl} = O - C_{-Alk} - X^{1} - N - Alk - X^{1} $

with R^1 , G, $-a^1=a^2-a^3=a^4$ -; Alk, X^1 R^2 and R^4 defined as in claim 1, in the presence of a suitable amination agent;

p) amination of an intermediate of formula (XIX)

$$H = C + C_{1-3} \text{alkyl} + NR^4$$

$$(XIX)$$

$$Q_6 N + CH_2 + C_{1-3} \text{alkyl} + NR^4$$

$$(XIX)$$

$$(XX)$$

$$(I-p)$$

with R^1 , G, and $-a^1=a^2-a^3=a^4$ defined as in claim 1, and $Q_6N-CH_2-C_{1-3}$ alkyl- NR^4

being defined as Q according to claim 1 provided that in the definition of Q, X^2 is C_{2-4} alkyl-NR⁴, in the presence of a suitable amination agent;

q) deprotecting an intermediate of formula (XXI)

$$P = O = G_1$$

$$Q = N$$

with R¹, Q, and -a¹=a²-a³=a⁴- defined as in claim 1, and HO-G₁ being defined as G according to claim 1 provided that G is substituted with hydroxy or HO-(CH₂CH₂O-)_n;

r) reducing an intermediate of formula (XXII)

- with R¹, Q, and -a¹=a²-a³=a⁴- defined as in claim 1, and H-G₂-OH being defined as G according to claim 1 provided that G is substituted with hydroxy and the carbon atom carrying the hydroxy substituent carries also at least one hydrogen, in the presence of a suitable reducing agent.
- and, if desired, converting compounds of formula (I) into each other following artknown transformations, and further, if desired, converting the compounds of
 formula (I), into a therapeutically active non-toxic acid addition salt by treatment
 with an acid, or into a therapeutically active non-toxic base addition salt by
 treatment with a base, or conversely, converting the acid addition salt form into the
 free base by treatment with alkali, or converting the base addition salt into the free
 acid by treatment with acid; and, if desired, preparing stereochemically isomeric
 forms, metal complexes, quaternary amines or N-oxide forms thereof.
- 14. A product containing (a) a compound as defined in claim 1, and (b) another
 25 antiviral compound, as a combined preparation for simultaneous, separate or sequential use in the treatment or the prevention of viral infections.

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15. A pharmaceutical composition comprising a pharmaceutically acceptable carrier and as active ingredients (a) a compound as defined in claim 1, and (b) another antiviral compound.

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B. FIELDS SE				<u></u>				
Minimum docu IPC 7	mentation searched (cl CO7D A61K	assification system follow A61P	ed by classification symb	pols)	•			
Documentation	n searched other than m	inimum documentation to	the extent that such doo	uments are included in t	the fields searched			
	•	the international search (where practical, search	terms used)			
	•	,						
C. DOCUMEN	ITS CONSIDERED TO E	BE RELEVANT						
Category ° (Citation of document, wit	th indication, where appro	opriate, of the relevant pa	issages	Relevant to claim No.			
A	6 February	' A (JANSSEN P 1992 (1992-02 ne 9 - line 1	-06)	NV)	1,7			
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Further	r documents are listed in	the continuation of box 0	: X	Patent family members	are listed in annex.			
° Special cateç	gories of cited documents	s:						
	defining the general stated to be of particular rele		or ₍	priority date and not in co	er the international filing date onflict with the application but ciple or theory underlying the			
filing date	•	or after the international	"X" docu car	iment of particular releva	ance; the claimed invention or cannot be considered to			
which is o	which may throw doubts cited to establish the pub ir other special reason (a	plication date of another s s specified)	"Y" docu car	ment of particular releva	nen the document is taken alone unce; the claimed invention olve an inventive step when the			
other mea "P" document	ans published prior to the int	osure, use, exhibition or temational filing date but	me in t	nts, such combination be he art.	one or more other such docu- sing obvious to a person skilled			
later than	the priority date claimed	1		ment member of the sar e of mailing of the interna	<u> </u>			
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	European Patent Omc NL - 2280 HV Rijswiji Tel. (+31-70) 340-204 Fax: (+31-70) 340-30	40, Tx. 31 651 epo nl,		Alfaro Faus,	I			

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A. CLASSIF IPC 7	CO7D491/056, 319:00, 221:00) (C07D491/056, 319:00, 221:00)	7D471/04,235:00,221:00),	
According to	International Patent Classification (IPC) or to both national class	ification and IPC	
B. FIELDS	SEARCHED		
Minimum do	currientation searched (classification system followed by classification s	cation symbols)	
Documentati	ion searched other than minimum documentation to the extent th	at such documents are included in the fields se	arched
	ata base consulted during the international search (name of data	base and, where practical, search terms used	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the	e relevant passages	Relevant to claim No.
Furt	her documents are listed in the continuation of box C.	Patent family members are listed	in annex.
"A" docum consist "E" earlier filing of "L" docum which citatic "O" docum other "P" docum later t	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or its cited to establish the publication date of another or or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but than the priority date claimed	T' later document published after the interest or priority date and not in conflict with cited to understand the principle or the invention 'X' document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the do 'Y' document of particular relevance; the cannot be considered to involve an indocument is combined with one or moments, such combination being obvious in the art. '&' document member of the same patent. 'Date of mailing of the international sea	the application but cory underlying the lairned invention be considered to current is taken alone lairned invention rentive step when the re other such docusto a person skilled
	mailing address of the ISA	Authorized officer	· · · · · · · · · · · · · · · · · · ·
Hanle all	European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Eav. (431-70) 340-3018	Alfaro Faus, I	

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FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Present claims 1 to 15. relate to an extremely large number of possible compounds. Support within the meaning of Article 6 PCT and/or disclosure within the meaning of Article 5 PCT is to be found, however, for only a very small proportion of the compounds claimed. In the present case, the claims so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible. Consequently, the search has been carried out for those parts of the claims which appear to be supported and disclosed, namely those parts relating to the compounds of formula I where Q is a 1-R2-piperidy1-4-amino or amino(cyclo)alkylamino group and their intermediates as described in the examples of tables 3 to 13.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

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Interna: il Application No PCT/EP 00/05677

Patent document ited in search report	ļ	Publication date		atent family nember(s)	Publication date
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(19) World Intellectual Property Organization
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- (21) International Application Number: PCT/EP00/05677
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- (25) Filing Language:

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(54) Title: RESPIRATORY SYNCYTIAL VIRUS REPLICATION INHIBITORS

(57) Abstract: The present invention concerns compounds of formula (I), prodrugs, N-oxides, addition salts, quaternary amines, metal complexes and stereochemically isomeric forms thereof wherein -a1=a2-a3=a4- represents a radical of formula -CH=CH-CH=CH-; -N=CH-CH=CH-; -CH=N-CH=CH-; -CH=CH-N=CH-; -CH=CH-CH=N-; wherein each hydrogen atom may optionally be substituted; Q is a radical of formulae (b-1), (b-2), (b-3), (b-4), (b-5), (b-6), (b-7) and (b-8), wherein Alk is C₁₋₆alkanediyl; Y¹ is a bivalent radical of formula -NR²- or -CH(NR²R⁴)-; X¹ is NR⁴, S, S(=O), S(=O)₂, O, CH₂, C(=O)₃ CH(=CH₂), CH(OH), CH(CH₃), CH(OCH₃), CH(SCH₃), CH(NR^{5a}R^{5b}), CH₂-NR⁴ or NR⁴-CH₂; X² is a direct bond, CH₂, C(=O), NR⁴, C₁₋₄alkyl-NR⁴, NR⁴-C₁₋₄alkyl, t is 2 to 5; u is 1 to 5; v is 2 or 3; and whereby each hydrogen in Alk and in (b-3), (b-4), (b-5), (b-6), (b-7) and (b-8), may optionally be replaced by R3; provided that when R3 is hydroxy or C1.6alkyloxy, then R3 cannot replace a hydrogen atom in the α position relative to a nitrogen atom; G is a direct bond or optionally substituted C_{1.00}alkanediyl; R¹ is an optionally substituted bicyclic heterocycle; R² is hydrogen, formyl, C_{1.6}alkylcarbonyl, Hetcarbonyl, pyrrolidinyl, piperidinyl, homopiperidinyl, C_{3-7} cycloalkyl or $C_{1.10}$ alkyl substituted with $N(R^6)_2$ and optionally with another substituent; R^3 is hydrogen, hydroxy, C_{1.6}alkyl, C_{1.6}alkyloxy, arylC_{1.6}alkyl or arylC_{1.6}alkyloxy, R⁴ is hydrogen, C_{1.6}alkyl or arylC_{1.6}alkyl; R^{5a}, R^{5b}, R^{5c} and R^{5d} are hydrogen or C₁₋₆alkyl; or R^{5a} and R^{5b}, or R^{5c} and R^{5d} taken together from a bivalent radical of formula -(CH₂)_s- wherein S is 4 or 5; R^6 is hydrogen, C_{1-6} alkyl, formyl, hydroxy C_{1-6} alkyl, C_{1-6} alkyl, aryl is optionally substituted phenyl; Het is pyridyl, pyrimidinyl, pyryzinyl, pyridazinyl; as respiratory syncytial virus replication inhibitors; their preparation, compositions containing them and their use as a medicine.



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